



S E R V I C E   G U I D E

# Avalon Fetal Monitor FM20 / FM30

F E T A L   M O N I T O R I N G

**PHILIPS**

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# Introduction

This Service Guide contains technical details for the Avalon FM20 and FM30 Fetal/Maternal Monitors. It provides a technical foundation to support effective troubleshooting and repair. It is not a comprehensive, in-depth explanation of the product architecture or technical implementation. It offers enough information on the functions and operations of the monitoring systems so that engineers who repair them are better able to understand how they work. It covers the physiological measurements and the monitor hardware that acquires and displays them.

The *Avalon FM20/FM30 Fetal Monitor Service Guide* supplements the maintenance and troubleshooting procedures, carried out by the operator, that are described in the *Instructions for Use*. Refer to the *Instructions for Use* for maintenance and troubleshooting procedures that may be performed during normal operation.

Only qualified service personnel should attempt to install the system, disassemble the monitor, remove or replace any internal assemblies, or replace the transducer cable or belt buttons.

## Who Should Read This Guide

This guide is for biomedical engineers or technicians responsible for troubleshooting, repairing, and maintaining Philips' Avalon fetal monitors.

You must:

- understand English
- be familiar with standard medical equipment installation procedures
- be familiar with current conventional technical terms as used throughout this guide

## What to Do Next

Familiarize yourself with the contents of this guide and the *Instructions for Use* before attempting to service or repair the system.

# Repair Strategy

The Service Support Tool software helps you to determine whether a fault is a hardware or software problem. The main replaceable parts are:

- unit exchange for the transducers
- replacement of
  - the top cover assembly
  - the bottom housing
  - the power supply assembly
  - the display assembly
  - the recorder adapter board
  - the paper drawer assembly
  - the paper sensor assembly
  - the stepper motor assembly
  - the SpO<sub>2</sub> assembly
  - the noninvasive blood pressure assembly
  - the main CPU board
  - the bus master board
  - the socket connector block
  - the transducer cable
  - the transducer belt button

See Chapter 9, “Parts” for part numbers, and Chapter 10, “Disassembly and Reassembly” for repair details.

Repair or replacement of individual components on the boards is not supported, and should never be attempted.

For tests that you are required to perform after repairs, refer to “When to Perform Test Blocks” on page 29.

# Manufacturer's Information

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Philips Medizin Systeme Böblingen GmbH  
Hewlett-Packard-Str. 2

71034 Böblingen, Germany

## Passwords

In order to access different modes within the monitor a password may be required. The passwords are listed below.

Monitoring Mode: No password required

Configuration Mode: 71034

Demo Mode: 14432

Service Mode: 1345

Refer to Chapter 12, “Understanding Configuration” before making any changes to the monitor configuration.

## Warnings and Cautions

In this guide:

- A **warning** alerts you to a potential serious outcome, adverse event or safety hazard. Failure to observe a warning may result in death or serious injury to the user or patient.
- A **caution** alerts you where special care is necessary for the safe and effective use of the product. Failure to observe a caution may result in minor or moderate personal injury or damage to the product or other property, and possibly in a remote risk of more serious injury.



# Site Preparation

## Introduction

This section describes the procedures you should follow to plan and prepare a site for an Avalon FM20/FM30 fetal monitor installation.

- Site planning.
- Roles and responsibilities for local and Philips personnel.

## Site Planning

The careful planning of the site for the FM20/FM30 monitor is essential for its safe and efficient operation. A consulting schedule should be established between the Customer and Philips Sales and Support Representatives, to ensure that all preparations are completed when the system is delivered.

The site planning phases prior to equipment installation are:

**Location:** Planning the location of the various system components.

**Environment:** Confirming and correcting, as necessary, the environment of the proposed installation site(s).

**System Capabilities:** Explaining the possibilities for system expansion.

**Mounting:** Referencing the mounting hardware information website for the listing of suitable mounting hardware recommended for use with the various system components, and all details on the available mounts and accessories.

**Cabling:** Identifying the requirements for the cabling, conduiting and faceplates for connecting the various system components.

## Roles and Responsibilities

This section describes the procedures necessary to prepare a site for a system installation. The procedures are grouped into two parts: procedures that local staff or contractors are responsible for, and procedures that Philips personnel are responsible for.

### Site Preparation Responsibilities

#### Local Staff

- Ensure that all safety, environmental and power requirements are met.
- Provide power outlets.
- Prepare mounts, and consult Philips for detailed mounting requirements.

- Pull cables, install conduit, install wallboxes.

#### Philips Personnel

- Provide the customer with the safety, environmental and power requirements.
- Assemble mounts, as necessary.
- Provide requirements for cabling.

### Procedures for Local Staff

The following tasks must be completed **before** the procedures for Philips personnel may be started.

- Providing Power Outlets

Provide a power outlet in the vicinity (1 m or 3 ft) or any peripheral equipment.

---

**WARNING** Always use the supplied power cord with the earthed mains plug to connect the monitor to an earthed AC mains socket. Never adapt the mains plug from the power supply to fit an unearthing AC mains socket.

---

- Preparing Mounts

Where ceiling, wall, or shelf mounts are required for mounting the equipment, the customer is responsible for the following:

- Providing and installing all hardware which is required to install the mounting hardware supplied by Philips as detailed in the installation notes.
- Making sure that all ceilings, walls, and mounting rails that supports mounting hardware are suitable for their proposed load.

---

**WARNING** It is the customer's responsibility to have the attachment of the mounting hardware to the ceiling, wall, or mounting rail and the construction of the ceiling, wall, or mounting rail evaluated for structural integrity and compliance with all local, state and any other required codes by a registered, professional, structural and/or mechanical engineer.

Although considerable effort has been made to ensure the safety of the ceiling mount installation and or mounting guidelines, it is to be understood that the installation itself is beyond the control of Philips Medical Systems. Accordingly, Philips Medical Systems will not be responsible for the failure of any such installation.

---

- Providing Conduit

- Providing conduit and/or trunking of a sufficient cross-sectional area for the planned cables and possible future expansion (for additional components or systems).
- Providing and/or installing suitable wall boxes to accommodate the faceplates.

- Pulling Cables

---

**WARNING** NEVER run power cables through the same conduit or trunking used for system cables.

---

- Installing Wall Boxes

It is the customer's responsibility to provide and install wallboxes to house faceplates. The customer must notify the Philips installation coordinator of which size is to be used.

## Procedures for Philips Personnel

Before you begin the procedures in the installation sections, ensure that the customer has completed all necessary preparations outlined in the previous section, "Procedures for Local Staff."

# Site Requirements

The site requirements are listed in this section.

## Space Requirements

The situating of the monitor should be planned such that the nursing staff are able to monitor the patient with relative ease, with all patient connectors and controls readily available and the displays clearly visible. The location should also allow access to service personnel without excessive disruption and should have sufficient clearance all round to allow air circulation.

Dimensions and weight:

**Monitor:**

Size (W x H x D): 335 x 286 x 133 mm (13.2 x 11.3 x 5.2 in)

Weight; 5.1 kg (11.2 lb)

**Transducer:**

Size (diameter): 83 mm (3.27 in)

Weight (without cable): 190g (6.7 oz.)

## Environmental Requirements

The environment where the FM20/FM30 monitor will be used should be reasonably free from vibration, dust and corrosive or explosive gases. The ambient operating and storage conditions for the FM20/FM30 monitor must be observed. If these conditions are not met, the accuracy of the system will be affected and damage can occur.

<b>Monitor (M2702A/M2703A); Interface Cable for Avalon CTS (M2731-60001)</b>		
<b>Temperature Range</b>	Operating	0°C to 45°C (32°F to 113°F)
	Storage	-20°C to 60°C (-4°F to 140°F)
<b>Humidity Range</b>	Operating	<95% relative humidity @ 40°C/104°F
	Storage	<90% relative humidity @ 60°C/140°F
<b>Altitude Range</b>	Operating	-500 to 3000 m/-1640 to 9840 ft.
	Storage	-500 to 13100 m/-1640 to 43000 ft.

<b>Transducers (M2734A/M2735A/M2736A/M2738A)</b>		
<b>Humidity Range</b>	Operating	<95% relative humidity @ 40°C/104°F
	Storage	<90% relative humidity @ 60°C/140°F
<b>Altitude Range</b>	Operating	-500 to 3000 m/-1640 to 9840 ft.
	Storage	-500 to 13100 m/-1640 to 43000 ft.

<b>SpO<sub>2</sub> Sensors</b>	
<b>Operating Temperature Range</b>	0°C to 37°C (32°F to 98.6°F)

## Safety Requirements (Customer or Philips)

The monitor is an electrical Class II device in which the protection against electric shock does not rely on basic insulation and a protective earth conductor but on double and/or reinforced insulation.

- 
- WARNING**
- Always use the supplied power cord with the earthed mains plug to connect the monitor to an earthed AC mains socket. Never adapt the mains plug from the power supply to fit an unearthed AC mains socket.
  - The protective earth conductor is required for EMC purposes. It has no protective function against electric shock! The protection against electric shock in this device is provided by double and/or reinforced insulation.
  - Do not use additional AC mains extension cords or multiple portable socket-outlets. If a multiple portable socket-outlet without an approved separating transformer is used, the interruption of its protective earthing may result in equipment leakage currents equal to the sum of the individual earth leakage currents, so exceeding allowable limits.
- 

## Electrical Requirements (Customer or Philips)

Line Voltage Connection

The FM20/FM30 monitor uses < 60 W.

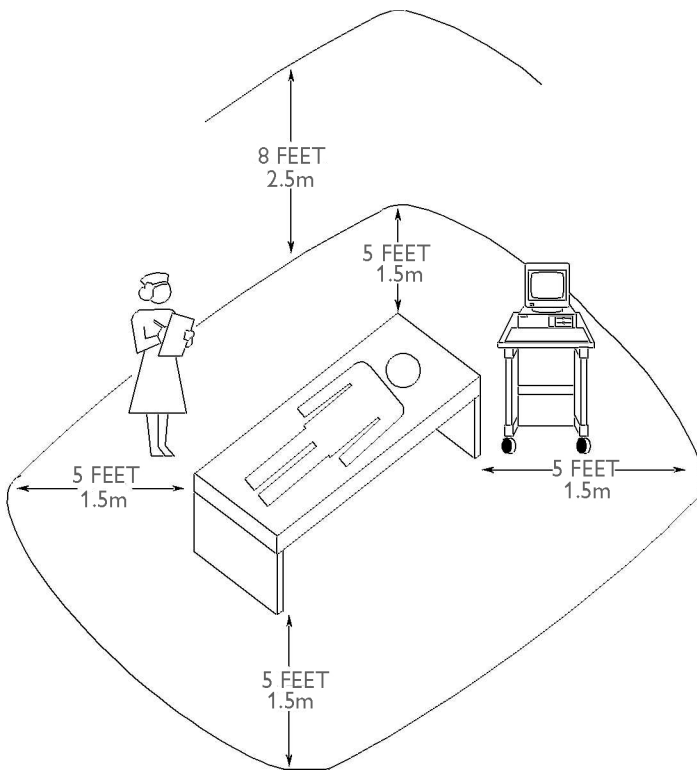
Line Voltage: the FM20/FM30 monitor may be operated on ac line voltage ranges of 100 to 240V (50/60 Hz).

## Connecting Non-Medical Devices

The standard IEC/EN 60601-1-1 applies to any combination of devices, where at least one is a medical electrical device. Therefore IEC/EN 60601-1-1 must still be met after all devices are connected.

- WARNING**
- Do not use a device in the patient vicinity if it does not comply with IEC/EN 60601-1. The whole installation, including devices outside of the patient vicinity, must comply with IEC/EN 60601-1-1. Any non-medical device, **including a PC running an OB TraceVue system**, placed and operated in the patient's vicinity must be powered via a separating transformer (compliant with IEC/EN 60601-1-1) that ensures mechanical fixing of the power cords and covering of any unused power outlets.
  - Do not connect any devices that are not supported as part of a system.

Whenever you combine equipment to form a system, for example, connecting the monitor to an OB TraceVue system, perform a system test according to IEC/EN 60601-1-1 (see "System Test" on page 42).



**Figure 1 Equipment Location in the Patient Vicinity**

## Cabling Options and Requirements for Connection to OB TraceVue

For cabling options and requirements for connection to an OB TraceVue system, refer to the *OB TraceVue Site Preparation Guide* and the *OB TraceVue Service Guide*.

## Mounting Options

See "Mounting Hardware" on page 91 for a list of mounting options. Refer to "Mounting Instructions" on page 12, or contact your local Philips representative for advice on mounting the monitor.

## Input Devices

The following table describes the input devices which can be connected to the monitor via the optional PS/2 interface.

Product Option Number	Part Number	12NC Part Number	Description
M8024A #A01	862454	9898 031 24741	Slimline Keyboard with integrated Trackball
M8024A #B01	M4046-60104	4512 610 00661	Optical Mouse USB / PS/2
M8024A #C01	M4046-60103	4512 610 00651	Wired Track Ball USB / PS2
M8024A #C02	M4046-60105	4512 610 00671	Wireless Track Ball
M8024A #C03	M4046-60106	4512 610 00681	Wired off table Track Mouse

# Installation Instructions

The information contained in this chapter, **in addition to that given in the *Instructions for Use***, should enable the monitor to be installed ready for use (the preparation and planning should be adhered to as specified in the “Site Preparation” chapter). Safety checks and inspection procedures for mounts are explained in the “Testing and Maintenance” chapter, and configuration of the system is explained in the “Configuration” chapter.

Please keep the packing materials until you have completed the initial inspection, in case there is a defect on arrival.

## Initial Inspection

Inspect the delivery on arrival.

### Visual Inspection

Open the shipping container(s) and examine each part of the instrument for visible damage, such as broken connectors or controls, or scratches on the equipment surfaces. If the shipping carton/container is undamaged, check the cushioning material and note any signs of severe stress as an indication of rough handling in transit. This may be necessary to support claims for hidden damage that may only become apparent during subsequent testing.

### Electrical Inspection

The instrument has undergone extensive testing prior to shipment. Safety testing at installation is not required (except in situations where devices are interconnected forming a system, see “Connecting Non-Medical Devices” on page 8). An extensive self check may be performed. This recommendation does not supersede local requirements.

All tests are described in the “Testing and Maintenance” chapter of this manual.

## Claims for Damage

When the equipment is received, if physical damage is evident or if the monitor does not meet the specified operational requirements of the patient safety checks or the extended self check, notify the carrier and the nearest Philips Sales/Support Office at once. Philips will arrange for immediate repair or replacement of the instrument without waiting for the claim settlement by the carrier.

## Rereading for Shipment or Storage

If the instrument is to be shipped to a Philips Sales/Support Office, securely attach a label showing the name and address of the owner, the instrument model and serial numbers, and the repair required (or symptoms of the fault). If available and reusable, the original Philips packaging should be used to provide adequate protection during transit. If the original Philips packaging is not available or reusable please contact the Philips Sales/Support Office who will provide information about adequate packaging materials and methods.

## Mounting Instructions

Every type of compatible mounting solution is delivered with a complete set of mounting hardware and instructions. Refer to the Site prep chapter for a list of mounting options. Refer to the documentation delivered with the mounting hardware for instructions on assembling mounts.

**WARNING** It is the customer's responsibility to have the attachment of the mounting hardware to the ceiling, wall, or mounting rail and the construction of the ceiling, wall, or mounting rail evaluated for structural integrity and compliance with all local, state and any other required codes by a registered, professional, structural and/or mechanical engineer.

Ensure that this commitment has been met before assembling mounts.

## Line Voltage Selection

You do not need to set the line voltage, as this is done automatically by the power supply. The monitor has a wide-range power supply that allows you to operate the monitor from an AC (alternating current) power source of 100 V to 240 V ( $\pm$  10%) and 50/60 Hz ( $\pm$  5%).

## Connecting the Monitor to AC Mains

The monitor is an electrical Class II device in which the protection against electric shock does not rely on basic insulation and a protective earth conductor but on double and/or reinforced insulation.

**WARNING**

- Always use the supplied power cord with the earthed mains plug to connect the monitor to an earthed AC mains socket. Never adapt the mains plug from the power supply to fit an unearthed AC mains socket.
- The protective earth conductor is required for EMC purposes. It has no protective function against electric shock! The protection against electric shock in this device is provided by double and/or reinforced insulation.
- Do not use AC mains extension cords or multiple portable socket-outlets.

# Connecting the Monitor to Non-Medical Devices

Connect the monitor to an obstetrical surveillance system, such as OB TraceVue, via the optional system interface. For cabling requirements, refer to “Cabling Options and Requirements for Connection to OB TraceVue” on page 9. For safety-related information, refer to “Connecting Non-Medical Devices” on page 8, and “System Test” on page 42.

## Before Using the Monitor

- 
- WARNING**
- Before starting monitoring, check that the configuration meets your requirements.
  - Be aware that the monitors in your care area may each have different alarm settings, to suit different patients. Always check that the alarm settings are appropriate for your patient before you start monitoring.
- 

Check that the following configuration settings are suitable:

- Line Frequency
- Paper Scale
- Paper Speed
- Equipment Label
- Configured SmartKeys
- Input device configuration (if using an external keyboard or mouse)

If you need to enter configuration mode to change settings:

- 1 In the **Main Setup** menu, select **Operating Modes**.
- 2 Select **Config** and enter the passcode.  
The passcode for configuration mode is given in the monitor’s service documentation.

The monitor displays **Config** at the right hand side of the status line and in the center of the Screen while you are in configuration mode.

Before you leave configuration mode, always be sure to store any changes you made. You must store changes made to each Settings Block and to each Profile, individually. As it may be difficult to remember whether the settings you changed belong to a Monitor Settings block or a Measurement Settings block, we recommend that you store each block before you leave configuration mode.

To leave configuration mode:

- 1 Enter the **Main Setup** menu.
- 2 Select **Operating Modes**.
- 3 Select **Monitoring**.

## Checking and Setting Line Frequency

Before using the monitor, check that the line frequency setting is correct for your location, and change the setting if necessary in Service Mode.

**WARNING** An incorrect line frequency setting can affect the ECG filter, and disturb the ECG measurement. Ensure the line frequency setting is correct.

To set the line frequency:

- 1 Enter the **Main Setup** menu.
- 2 Select **Global Settings**.
- 3 Select **Line Frequency** and select **50Hz** or **60Hz** from the pop-up list.

## Checking/Setting Paper Scale

Check the paper Scale Type (**US** for paper with a scale of 30-240, or **Internat' 1** for paper with a scale of 50-210) in the Fetal Recorder menu. In Monitoring Mode, you can see this setting (grayed out), but you cannot change it. It can be changed in Configuration Mode.

- 1 Enter the **Main Setup** menu by selecting the SmartKey  .
- 2 Select **Fetal Recorder**.
- 3 Check the current setting for Scale Type. If it is not appropriate, change it in the Fetal Recorder menu in Configuration Mode:  
Select **Scale Type** to toggle between **US** and **Internat' 1**.

## Checking/Setting Paper Speed

Check the paper speed before using the monitor. You can choose a paper speed of 1, 2, or 3 centimeters per minute (cm/min). The default setting is 3 cm/min. In Monitoring Mode, you can see this setting (grayed out), but you cannot change it. It can be changed in Configuration Mode.

As a change in paper speed results in a change in the appearance of a FHR trace, you are advised to ensure **ALL** monitors in your institution are set to the same speed.

To set the paper speed:

- 1 Enter the **Main Setup** menu using the SmartKey  .
- 2 Select **Fetal Recorder**.
- 3 In the Recorder menu, you can see the current speed setting. Select **Recorder Speed**.
- 4 Select the desired speed from the given choices: **1**, **2** or **3 cm/min**.

## Configuring the Equipment Label

OB TraceVue requires a unique equipment label. In OB TraceVue, it is possible to prevent connection to monitors with specific equipment labels by means of a filtering mechanism. For more details, see the OB TraceVue Instructions for Use.

- 1 Select the **Bed Label** screen element to call up the **Bed Info** menu.
- 2 Select **Equipment Label** to call up the onscreen keyboard.
- 3 Enter the system identifier.

## Configuring SmartKeys

Check that the configured SmartKeys are suitable. Configure the SmartKeys preferred by the institution from a global list Global Smart Keys. The global list of SmartKeys is stored as a unique monitor setting in the monitor configuration. See the section “Configuring Global SmartKeys” on page 180 for details on how to configure the global SmartKey list.

## PS/2 Keyboard/Mouse

Switch off the monitor before connecting any PS/2 compatible device.

Connect the PS/2 connector to the PS/2 Interface board in the monitor at the slot indicated by the appropriate symbol.

The default keyboard language setting for all initial configurations is “US”.

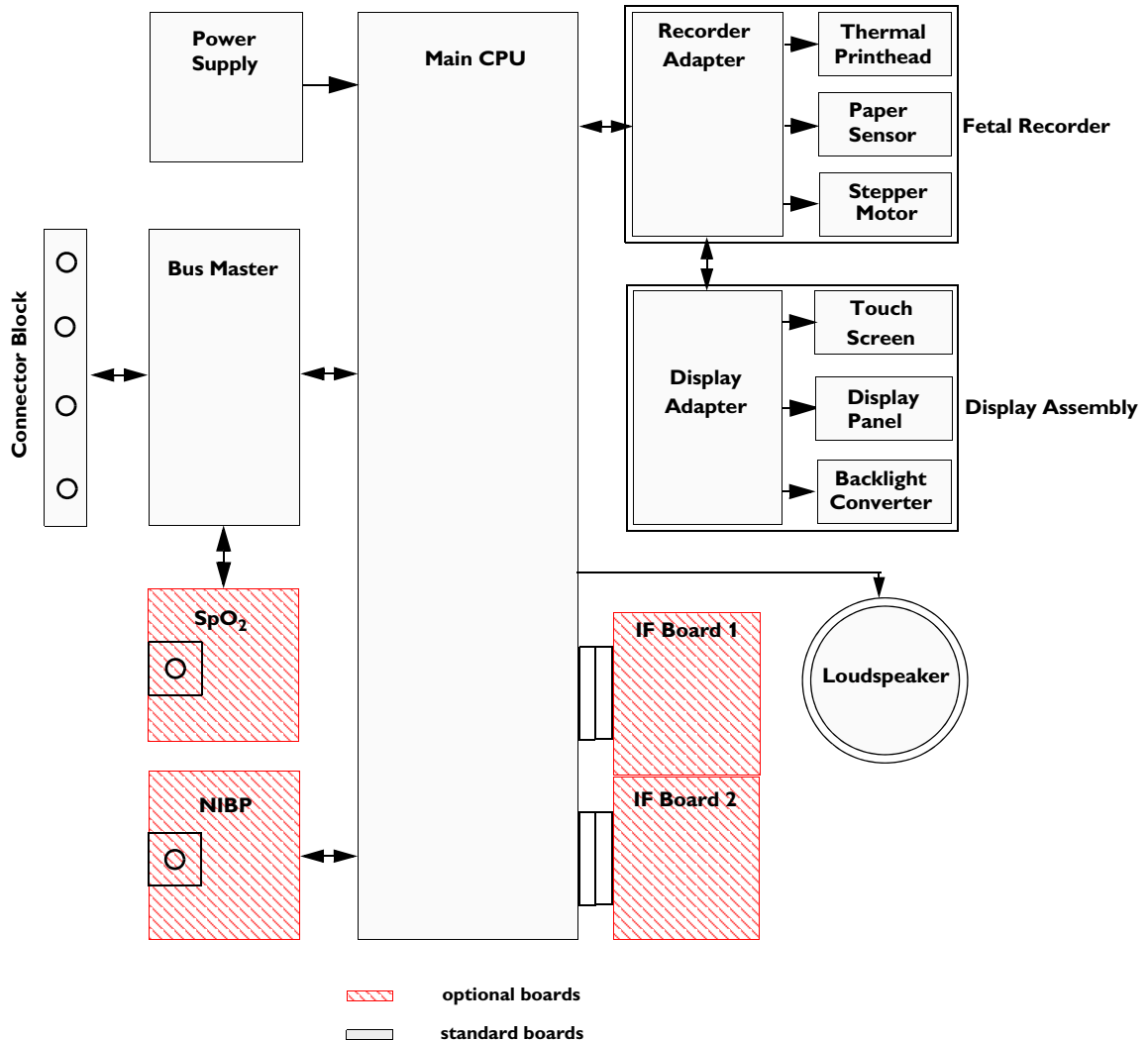
To configure the keyboard language manually, go to Service Mode, select **Main Setup -> Hardware -> Keyboard** and then select the proper language. Please note that this setting does not clone.



# Theory of Operation

This chapter describes the functional operation of the monitor and the transducers. It incorporates features of the mechanical design, indicating the physical relationship of the assemblies and components.

## Monitor Hardware Overview



The monitor consists of nine main functional components:

- Power supply M2703-60001
- Connector Block (1253-8415)
- Bus Master Board (M2703-66460)
- Main CPU Board (M2705-66510)
- Fetal Recorder (Thermal Printer Unit)
- Display Assembly (M2703-64503)
- Noninvasive Blood Pressure Board (optional, M2703-64502)
- SpO<sub>2</sub> Board (optional, M2703-66453)
- Input /Output Interface Boards (optional):
  - LAN / RS232 (M2703-67501)
  - Dual PS/2 (M8086-67501)

## Power Supply

The power supply is a wide-range input switching unit, with an output of 24V. It is located in the bottom housing assembly.

## Connector Block

Any compatible fetal transducer can be connected in any order to the monitor via the sockets on the Connector Block. The Connector Block is located on the Bus Master Board, and is exchangeable. The assembly made up of the Connector Block and the Bus Master Board is referred to as the Front End Assembly in the chapters “Disassembly and Reassembly” and “Upgrades”.

## Bus Master Board

The signals from the transducers or sensors are conveyed from the sensor sockets on the Connector Carrier Board (M2703-66421) to the Bus Master Board (M2703-66420).

The Bus Master Board is responsible for transducer detection, communicates with the connected transducers via a CAN bus, and communicates parameter data to the Main CPU Board via a serial link for further processing and display.

## Main CPU Board

The Main CPU Board controls the monitor’s human interface, and is responsible for the final processing of data from the Bus Master Board. It sends this data to the TFT display, and to the thermal printer unit for recording traces and other patient data. It also controls the optional LAN/RS232 and PS/2 interface boards.

# Fetal Recorder (Thermal Printer Unit)

The fetal recorder is located in the Top Cover Assembly. The recorder consists of the following major parts:

- Recorder Adapter Board
- Thermal Line Printhead (TLPH)
- Paper Sensor
- Stepper Motor

## Recorder Adapter Board

Recorder signals are handled by the Recorder Adapter Board (M2703-66430), connected to the Main CPU Board. Video signals to the display are also wired through this board, and connection to the Display Adapter Board is made via a silver-colored 50-pin ribbon cable.

The recorder unit, including the TLPH, is connected to the Recorder Adapter Board via a white 50-pin, ribbon cable. The stepper motor and the paper sensor are also connected to the Recorder Adapter Board. The Recorder Adapter Board is connected to the Main CPU Board via a 154-pin connector.

## Thermal Line Printhead (TLPH)

The TLPH is located on its own holder in the recorder chassis.

## Paper Sensor

The paper sensor hardware consists of a reflective light sensor that detects the black marks on the trace paper, and paper-out. It is attached to the RFI Bracket, and connected to the Recorder Adapter Board via a removable cable connector.

## Stepper Motor

The stepper motor is a bipolar motor controlled by a micro-stepping motor driver on the Recorder Adapter Board. The motor is located on the recorder chassis and is connected to the Recorder Adapter Board via a removable cable connector.

# LCD Display and Touchscreen

The LCD Display Assembly consists of a four-wire resistive touchscreen, a 6.5" TFT panel, and a backlight inverter, all connected to the Display Adapter Board (M2703-66440) and fitted into the display housing.

The board is connected to the Recorder Adapter Board (M2703-66430) via a 50-pin ribbon cable. The green power LED is incorporated into the Display Adapter Board.

# Noninvasive Blood Pressure Assembly

The optional Noninvasive Blood Pressure Assembly (M2703-64602) is located in the front lefthand corner of the bottom housing assembly. It is connected via a serial link to the Main CPU Board.

# SpO<sub>2</sub> Assembly

The optional SpO<sub>2</sub> Assembly (M2703-64603) is physically located on the Bus Master Board, but sends data directly to the Main CPU Board via a serial link.

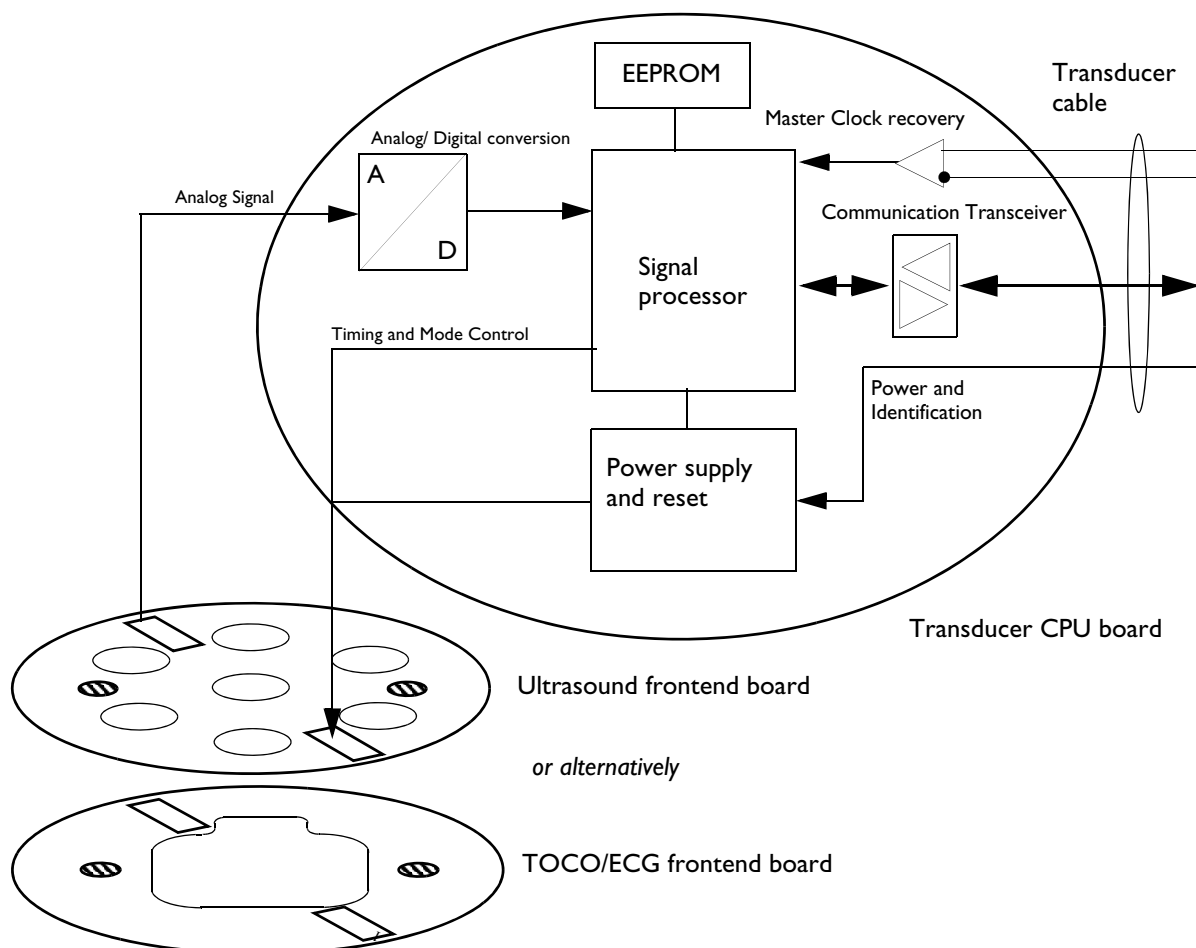
## Input/Output Interface Boards

There are two optional interface boards available:

- LAN/RS232 Interface Board, used for connecting to a PC running the Support Tool and to a surveillance and documentation system such OB TraceVue.
- PS/2 Interface Board, used for connecting an external keyboard or mouse.

The interface boards plug into the two interface slots on the underside of the device, and are controlled by the Main CPU Board.

## Transducer Hardware Overview



# Transducer Types

All transducers that can connect to the fetal sensor sockets can be used.

## Functional Description of the Transducer CPU

The CPU section of the transducers is made up of the following main functional blocks:

- CPU (micro controller)
- Analog-to-Digital Converter
- Communication Transceiver (CAN bus driver)
- EEPROM

### CPU (Micro Controller)

A single-chip processor is used to control the transducer, generate the frontend control signals, control the analog-to-digital signal conversion, and to perform the signal processing.

### Analog-to-Digital Converter

Analog-to-digital (A/D) signal conversion is carried out by the 16-bit AD converter. Digital signals are directly communicated from the A/D converter to the CPU.

### Communication Transceiver (CAN Bus Driver)

The communications transceiver (CAN bus driver) communicates directly with the transducer CPU, and allows the transducer to communicate with the Bus Master Board via the CAN bus.

### EEPROM

The serial EEPROM stores all non-volatile data required to operate the transducer (for example, calibration and correction factors for frontend gains and offsets, country-specific information, serial numbers and error logs).

## Toco Transducer Frontend

Uterine activity is measured by evaluating the hardness of the mother's abdomen with a pressure sensitive resistor bridge (strain gauge sensor element). The strain gauge sensor element requires an excitation voltage and its differential output signal is proportional to the pressure applied to it. A DC excitation voltage is used, and the resulting output signal is fed directly to an A/D signal converter before being sent to the processor.

## Ultrasound Transducer Frontend

The ultrasound frontend is a pulsed Doppler system with a 1.0 MHz ultrasound frequency, and a pulse repetition rate of 3 kHz. Seven ultrasound crystals are used as transmitter and receiver.

# Toco<sup>+</sup> Transducer Frontends

Several parameter frontends are combined on one board. In addition to the Toco frontend, additional supported parameters are DECG, MECG and IUP.

A seven-pin ‘D-type’ socket carries all parameter related inputs and outputs. An external mode resistor, connected to one of the pins, automatically detects which mode to set when an adapter cable is plugged in (whether it is DECG, MECG, or IUP).

## Toco Frontend

See “Toco Transducer Frontend” on page 21.

## IUP Frontend

Intrauterine pressure (IUP) is measured via a piezo resistive bridge with AC excitation connected to the RA / LA input pins of the ECG amplifier. A/D conversion of the IUP signal is done by the 16-bit A/D converter.

## ECG Frontend

The ECG frontend measures both DECG and MECG, using a 3-lead system (RA, LA and reference electrode). The ECG mode is automatically detected when an adapter cable is attached. Input lines are ESD protected.

# Patient Module Frontends

The patient module shares the same parameter frontends as the Toco<sup>+</sup> transducer, with the exception of the Toco frontend.

# Avalon CTS Interface Cable (TMIF)

The Avalon CTS Interface Cable contains the Telemetry Module Interface (TMIF). The TMIF shares the signal processing circuitry with the rest of the Avalon transducers, with the exception that it has no frontend board. The TMIF is responsible for converting the analog signals from the parameter frontends of the Avalon CTS transducers into digital signals for transmission to the fetal monitor.

# Interfaces

There are two interface boards available as options for the Avalon fetal monitors:

- LAN / RS232 system interface
- Dual PS/2 interface



The interfaces are “plug-and-play” boards, and fit into dedicated slots on the underside of the monitor. See “Removing the Interface Boards” on page 141 for details of how to remove and fit the boards.

## LAN / RS232 Interface

The LAN / RS232 system interface has two fully isolated ports:

- The LAN connection can be used for connecting the monitor to a PC for configuration or upgrade using the Support Tool, for connecting the monitor to an OB TraceVue obstetrical information system on a network, and for future system expansion.
- The RS232 connection can be used for connecting the monitor to an obstetrical information and surveillance system, such as OB TraceVue.

## Dual PS/2 Interface

This interface provides two PS/2 ports to enable the monitor to be connected to off-the-shelf, “plug-and-play” input devices:

- **Mouse:** any specified PS/2 mouse or trackball may be used for navigation and data entry.
- **Computer keyboard:** a PS/2 computer keyboard can be used for data entry instead of the on-screen pop-up keyboard.



# Connection to a Network

You can connect the fetal monitor to an OB TraceVue obstetrical information and surveillance system on a network using the LAN connection on the optional LAN / RS232 interface (see “LAN / RS232 Interface” on page 23).

## Network Infrastructure Requirements

The Avalon FM20/30 sends Connection Indication messages that OB TraceVue processes to establish an ethernet connection to the fetal monitor. The general requirements for connecting the Avalon FM20/30 to an OB TraceVue obstetrical surveillance system over a network are as follows:

- The fetal monitor and the data acquisition PC must be in the same network segment.
- The Avalon FM20/30 can be allocated a valid IP address either manually, or from a BOOTP service. If no IP address is entered manually, the Avalon FM20/30 requires a BOOTP service to obtain a valid IP address automatically, therefore BOOTP service must be available in each network segment. If the OB TraceVue internal server is part of that network segment, then it can be configured to serve BOOTP requests.
- The ethernet port of the Avalon FM20/30 supports only 10 M-bit half-duplex data transfer.

## Connection Indication Messages

Connection Indication (CI) messages can be sent by the Avalon FM20/30 in two ways: by Broadcast or by Unicast.

### Broadcast

When CI messages are sent by broadcast, they have the potential to reach any data acquisition PC in the same network segment, and the connection to the fetal monitor is made dynamically by the next available host PC (the data acquisition PC with the least active connections).

Broadcast is the default, and recommended, method for sending CI messages. This is because if there are multiple host PCs available in the same network segment, the broadcast method provides greater availability by allowing load balancing and failure-tolerant functionality. If a particular host PC happens to be unavailable, the next available PC takes over the connection. In the **Bed Information** menu, the **IP OB Server** entry is **0 . 0 . 0 . 0**.

## Unicast

When CI messages are sent by unicast, the fetal monitor sends a request to a specific target OB TraceVue data acquisition hosting PC in the same network segment. The CI message contains the IP address of the target PC, and only this PC will host the connection.

An example where CI messages are typically sent by unicast is where the fetal monitor and OB TraceVue PC are installed in the same cart, and you therefore always want the fetal monitor to be hosted by the same PC.

To avoid conflicts where there are multiple OB TraceVue systems operating in the same network segment, we recommend that you configure the fetal monitors to send the CI messages by unicast.

To enter the IP address of the target PC:

- 1 Enter Service Mode.
- 2 Select **Main Setup**.
- 3 Select **Bed Information**.
- 4 In the Bed Information menu, select **IP OB Server**.
- 5 Using the pop-up keypad, enter the IP address of the target server, and press **Enter** when you are done.

## Equipment Label and OB TraceVue Fetal Monitor Domain Name

For connection to an OB TraceVue system over a network, OB TraceVue requires each fetal monitor to have an unique equipment label.

When a fetal monitor is configured with an equipment label, this equipment label is sent as part of the CI message. In OB TraceVue, it is possible to specify a Fetal Monitor Domain Name in the fetal monitor configuration user interface (see OB TraceVue Instructions for Use for details). The OB TraceVue system compares this name with the fetal monitor equipment label. Only if the domain name matches the beginning of the equipment label string is the CI message processed and accepted. Otherwise the CI message is ignored.

Using this filtering process, you can avoid conflicts where there are multiple OB TraceVue systems operating in the same network segment by controlling which monitors connect to a specific OB TraceVue system, and which monitors are excluded.

To enter or change the equipment label:

- 1 Enter Configuration Mode.
- 2 Select **Main Setup**.
- 3 Select **Bed Information**.
- 4 In the Bed Information menu, select **Equipment Label**.
- 5 Enter the desired equipment label for your monitor (up to 12 alphanumeric characters are displayed), then press **Enter**.

# Testing and Maintenance

## Introduction

This chapter provides a checklist of the testing and maintenance procedures to ensure the performance and safety of the monitor and accessories.

These tests must be performed only by qualified personnel certified by the responsible organization. Qualifications required are: training on the subject, knowledge, experience and acquaintance with the relevant technologies, standards and local regulations. The personnel assessing safety must be able to recognize possible consequences and risks arising from non-conforming equipment.

All recurring safety and performance assurance tests must be performed under equal environmental conditions to be comparable.

Preventive Maintenance refers specifically to the series of tests required to make sure the measurement results are accurate. The accuracy and performance procedures are designed to be completed as specified in the following sections or when readings are in question.

For detailed instructions on the maintenance and cleaning of the monitor and its accessories, see Care and Cleaning, Maintenance and Troubleshooting in the monitor's Instructions for Use.

## Terminology and Definitions

The following terms and definitions are used throughout this chapter and taken from the international standards IEC 60601-1, IEC 60601-1-1 and IEC 62353.

**Medical System:** a medical electrical system is a combination of at least one medical electrical device and other electrical equipment, interconnected by functional connection or use of a multiple portable socket-outlet.

**Patient Vicinity:** any area in which intentional or unintentional contact can occur between the patient and parts of the medical system or between the patient and other persons who have had contact with parts of the medical system. The patient vicinity is defined anywhere within 1.5m (5 feet) of the perimeter of the patient's bed and 2.5m (8.2 feet) from the floor.

**Separation Device/Transformer:** a component or arrangement of components with input parts and output parts that, for safety reasons, prevent a transfer of unwanted voltage or current between parts of a medical system.

**Multiple Portable Socket-Outlet:** a combination of two or more socket-outlets intended to be connected to or integrated with flexible cables or cords, which can easily be moved from one place to another while connected to the power mains.

**Functional Connection:** an electrical connection for transfer of signals and/or power.

**Tests:** Safety or Performance Assurance test procedures which may consist of several steps.

## Recommended Frequency

Perform the procedures as indicated in the suggested testing timetable. These timetable recommendations do not supersede local requirements..

**Table 1: Suggested Testing Timetable**

Tests		Frequency
<b>Preventive Maintenance</b>	Noninvasive Blood Pressure Calibration	Once every two years, or as specified by local laws (whichever comes first).
<b>Other Regular Tests</b>	Visual Inspection	Before each use.
	Recorder Maintenance	Once a year, or if the printout is degraded.
	Testing Transducers and Patient Modules	Once a year, or if you suspect the measurement is incorrect.
<b>Performance Assurance Tests</b>	Noninvasive Blood Pressure Performance Tests	Once every two years, or if you suspect the measurement is incorrect.
	SpO <sub>2</sub> Performance	
<b>Safety Tests</b>	<b>Visual</b>	Visual Inspection After each service event.
	<b>Electrical</b>	Protective Earth Equipment Leakage Current Patient Leakage Current Once every two years and after repairs where the power supply is removed or replaced, or if the NBP assembly, SpO <sub>2</sub> board, or fetal sensor connector block is removed or replaced, or the monitor has been damaged by impact.
		System Test Once every two years.

## When to Perform Test Blocks

This table tells you when to perform specific test blocks. See page 57 for test details.

**Table 2: When to perform test blocks**

Service Event	Test Block(s) Required - Complete these tests
<b>Installation</b>	
Installation of standalone monitor	Perform Visual Inspection and Power On Tests
Installation of a monitor in combination with a medical or non-medical device connected to the same multiple socket outlet.	Perform Visual Inspection, Power On and System Tests
Installation of networked monitor (LAN)	Perform Visual Inspection, Power On and System tests
<b>Preventive Maintenance</b>	
Noninvasive Blood Pressure performance testing	Perform Noninvasive Blood Pressure Performance Test
<b>Other Regular Tests and Tasks</b>	
Visual Inspection	Perform Visual Inspection
Power On Test	Perform Power On Test
Transducer and Patient Module Testing	See “Transducers and Patient Modules; Functional Tests” on page 59.
Recorder Maintenance	Regular cleaning and maintenance (see “Transducers and Patient Modules; Functional Tests” on page 59) Perform the recorder selftest (see “Fetal Recorder Selftest Report” on page 66).
<b>Repairs</b>	
Repairs when the monitor has been damaged by impact, liquid ingressions, fire, short circuit or electrical surge.	Perform Visual Inspection, Power On, all Safety Tests and Full Performance Assurance Tests
Repairs where the power supply, the mains socket or an interface board is removed or replaced or the protective earth ground connection is disrupted.	Perform Visual Inspection, Power On, all Safety Tests and Basic Performance Assurance Tests
Repairs where the NBP pump has been replaced.	Perform Visual Inspection, Power On Test, all Safety Tests, Basic Performance Assurance Test and NBP Performance Test and Calibration.
All repair events involving any of the following: Noninvasive Blood Pressure connector, SpO <sub>2</sub> connector/board or the red fetal sensor sockets.	Perform Visual Inspection, Power On, Full Performance tests and all Safety Tests.
<b>Upgrades</b>	
Software Upgrades.	Perform Visual Inspection, Power On Test and Basic Performance Assurance Test unless otherwise specified in the Upgrade Installation Notes shipped with the upgrade.
Hardware Upgrades	Perform Visual Inspection, Power On Test and Basic Performance Assurance Test unless otherwise specified in the Upgrade Installation Notes shipped with the upgrade.

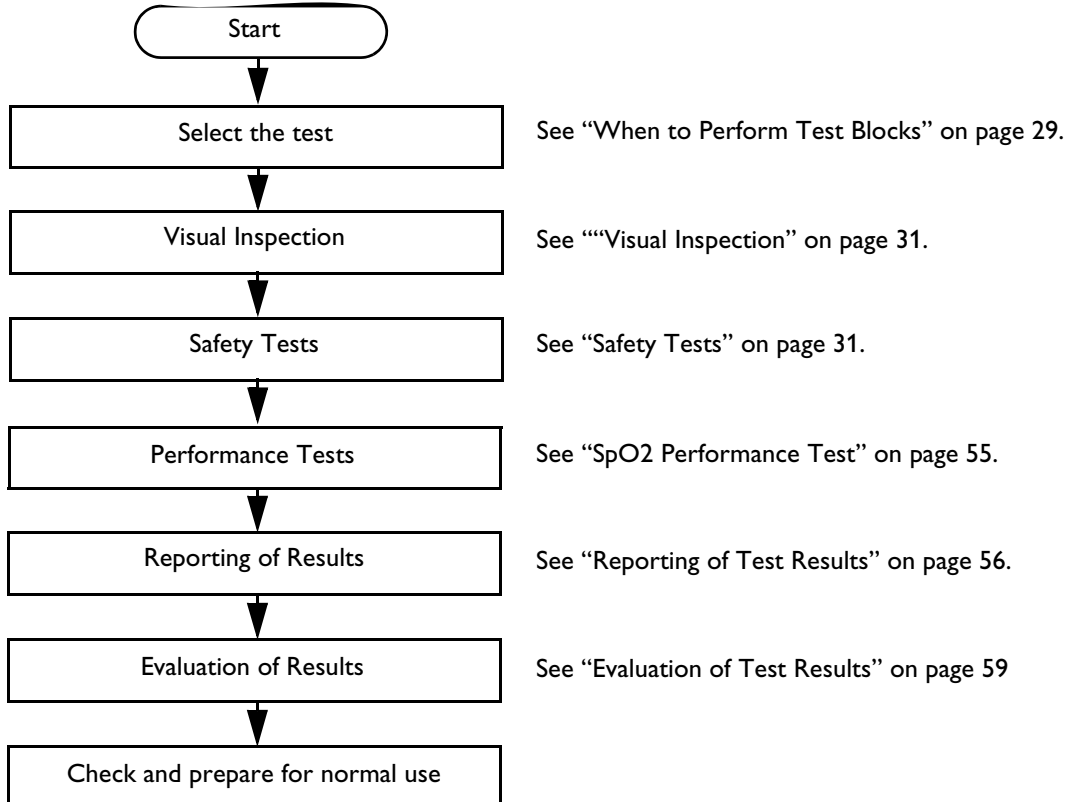
**Table 2: When to perform test blocks**

Service Event	Test Block(s) Required - Complete these tests
Installation of Interfaces or Hardware where the power supply or parameter boards need to be removed.	Perform Visual Inspection, Power On Test, Basic Performance Tests and all Safety Tests
Noninvasive Blood Pressure hardware upgrade M2702AU OptionB71.	Perform Visual Inspection, Power On, Noninvasive Blood Pressure Performance and all Safety tests (see Table 3).
SpO <sub>2</sub> hardware upgrade M2703AU Option B72.	Perform Visual Inspection, Power On, SpO <sub>2</sub> Performance and all Safety tests (see Table 3).
Noninvasive Blood Pressure/SpO <sub>2</sub> hardware upgrade M2703AU Option B73.	Perform Visual Inspection, Power On, Noninvasive Blood Pressure/SpO <sub>2</sub> Performance and all Safety tests (see Table 3).
<b>Combining or Exchanging System Components</b>	Perform the System Test (see Table 3 and “Touchscreen Calibration” on page 64).

**NOTE** It is the responsibility of the facility operator or their designee to obtain reference values for recurring safety and system tests. These reference values are the results of the first test cycles after an installation. You may also purchase this service from Philips.

## Testing Sequence

Here is a summary of the recommended sequence of testing:



**NOTE** If any single test fails, testing must be discontinued immediately and the device under test must be repaired or labeled as defective.

## Visual Inspection

### Before Each Use

Check all exterior housings for cracks and damage. Check the condition of all external cables, especially for splits or cracks and signs of twisting. If serious damage is evident, the cable should be replaced immediately. On the Toco<sup>+</sup> transducer and the patient module, ensure that the adapter cable socket is not damaged. Check that all mountings are correctly installed and secure. Refer to the instructions that accompany the relevant mounting solution.

### After Each Service, Maintenance or Repair Event

Ensure all fuses accessible from the outside comply with the manufacturer's specification.

Check:

- the integrity of mechanical parts, internally and externally.
- any damage or contamination, internally and externally.
- that no loose parts or foreign bodies remain in the device after servicing or repair.
- the integrity of all relevant accessories.

### Power On Test

- 1 Connect the monitoring system to mains and switch on the monitor.
- 2 Make sure that the green LED lights up and switches off again and the display then comes up.

The expected test result is pass: the monitor boots up and display comes up.

## Safety Tests

Safety tests are comprised of the following tests performed on the monitoring system:

- detachable power cord protective earth resistance (optional)
- equipment leakage current
- applied part leakage current
- system test (if required)

Safety test requirements are set according to international standards, their national deviations and specific local requirements. The safety tests detailed in this Service Guide are derived from international standards but may not be sufficient to meet local requirements. We recommend that you file the results of safety tests. This may help to identify a problem early particularly if the test results deteriorate over a period of time.

Each individual piece of equipment of the monitoring system which has its own connection to mains or which can be connected or disconnected from mains without the use of a tool must be tested individually. The monitoring system as a whole must be tested according to the System Test procedure.

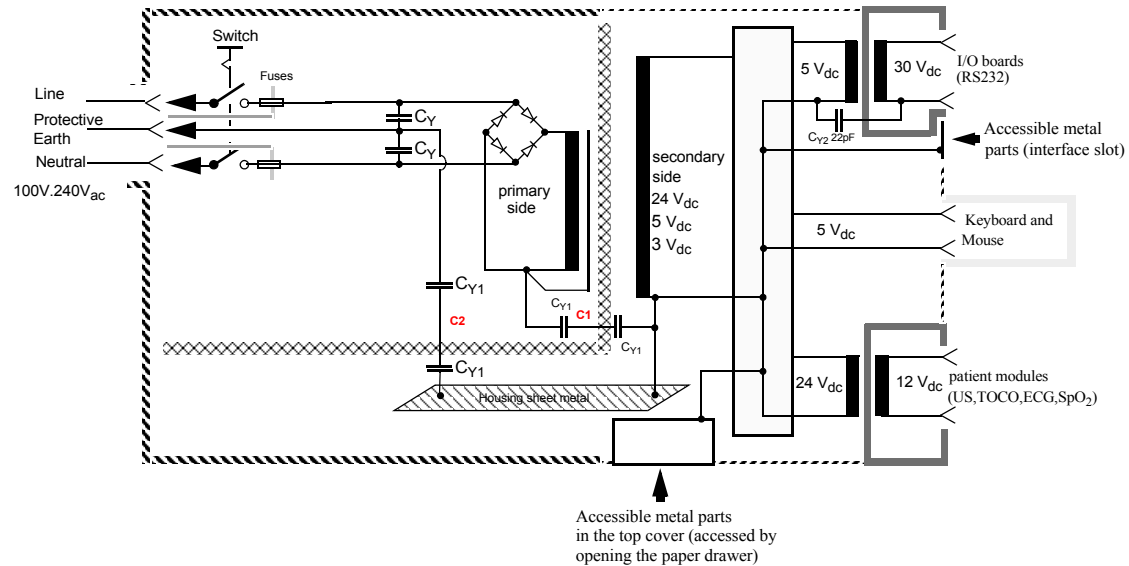
Accessories of the monitoring system which can affect the safety of the equipment under test or the results of the safety test must be included in the tests and documented.

## Warnings, Cautions, and Safety Precautions

- These tests are well established procedures of detecting abnormalities that, if undetected, could result in danger to either the patient or the operator.
- Disconnect the device under test from the patient before performing safety tests.
- Disconnect the device under test from mains before performing safety tests. If this is not possible, ensure that the performance of these tests does not result in danger to the safety analyzer operator, patients or other individuals.
- Test equipment (for example, a Safety Analyzer) is required to perform the safety tests. Please refer to Annex C of IEC/EN 62353 for exact requirements for the measurement equipment and for measurement circuits for protective earth resistance and leakage currents. Refer to the documentation that accompanies the test equipment. Only skilled technicians should perform safety testing.
- The consistent use of a *Safety Analyzer* as a routine step in closing a repair or upgrade is emphasized as a mandatory step to maintain approval agency status. You can also use the *Safety Analyzer* as a troubleshooting tool to detect abnormalities of line voltage and grounding plus total current loads.
- During safety testing, mains voltage and electrical currents are applied to the device under test. Ensure that there are no open electrical conductive parts during the performance of these tests. Avoid that users, patients or other individuals come into contact with touch voltage.
- For Europe and Asia/Pacific, the monitor complies with:  
IEC60601-1:1988 + A1:1991 + A2:1995 = EN60601-1:1990 +A1:1993 + A2:1995  
For USA, the monitor complies with:  
UL60601-1  
For Canada, CAN/CSA C22.2#601.1-M90
- Local regulations supersede the testing requirements listed in this chapter.
- If a non-medical device is connected to the medical electrical device during an installation, the resulting medical electrical system must comply with IEC/EN 60601-1-1.
- Perform safety tests as described on the following pages.

## Electrical Isolation Diagram

This diagram gives an overview of the electrical isolation of the monitor. Accessible metal parts are identified (see arrows).

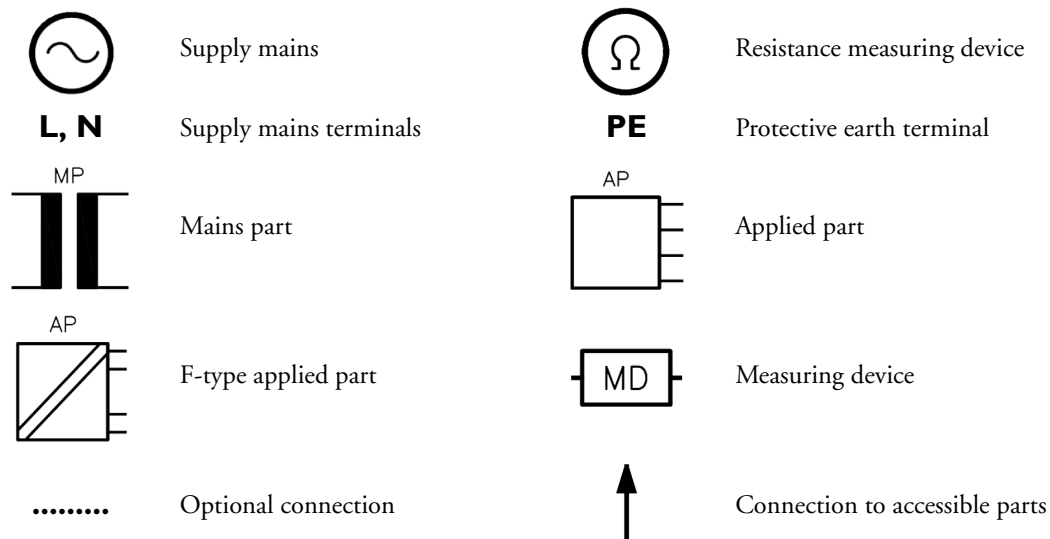


## Safety Test Procedures

Use the test procedures outlined here **only** for verifying safe installation or service of the product. The setups used for these tests and the acceptable ranges of values are derived from local and international standards but may not be equivalent. These tests are not a substitute for local safety testing where it is required for an installation or a service event. If using an approved safety tester, perform the tests in accordance with the information provided by the manufacturer of the tester and in accordance with your local regulations, for example IEC/EN 60601-1, UL60601-1 (US), IEC/EN 62353, and IEC/EN 60601-1-1. The safety tester should print results as detailed in this chapter, together with other data.

Please refer to Annex C of IEC/EN 62353 for requirements for the measurement equipment and for measurement circuits for protective earth resistance and leakage currents.

The following symbols are used in the diagrams illustrating the safety tests:



**CAUTION** After each service, maintenance or repair event:

Ensure all fuses accessible from the outside comply with the manufacturer's specification.

Check:

- the integrity of mechanical parts, internally and externally.
- any damage or contamination, internally and externally.
- that no loose parts or foreign bodies remain in the device after servicing or repair.
- the integrity of all relevant accessories.

**Hints for Correct Performance of Safety Tests**

- Perform a visual inspection on all detachable power cords used with the monitoring system and include these in all safety test procedures.
- Connection lines such as data lines or functional earth conductors may appear to act like protective earth connections. These may lead to incorrect measurements and need to be considered during testing. If necessary, unplug these connections.
- Position all cables and cords in such a manner that they do not influence the safety tests.
- Measurement of insulation resistance is not required.

## Guideline for the Performance of Safety Tests

Connect the detachable power cord of the device under test to the safety analyzer's test mains port. Connect the enclosure test lead of the safety analyzer to the enclosure of the device under test, e.g. to the equipotential connector. For testing the applied part leakage current, connect all applied parts to the safety analyzer using the appropriate patient lead or adapter cable. If necessary, repeat the safety test procedure until all available applied parts have been tested. Refer to the documentation that accompanies the safety analyzer for further details on how to set up and perform the test. Power on the device under test for all tests.

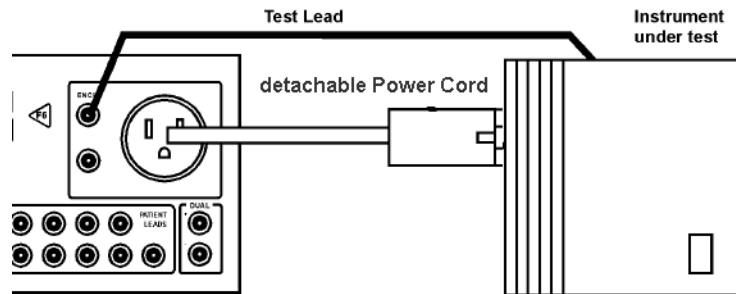


Figure 2 Protective Earth Resistance Test - Setup Example

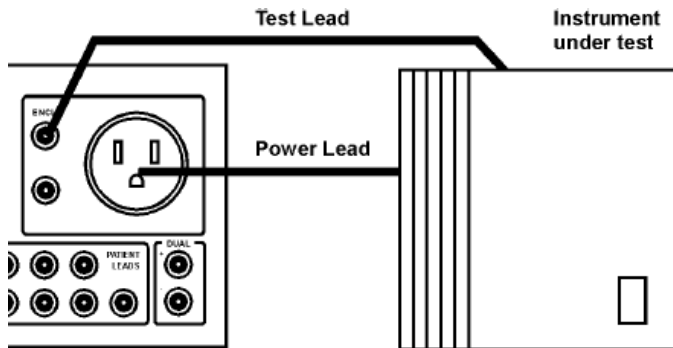
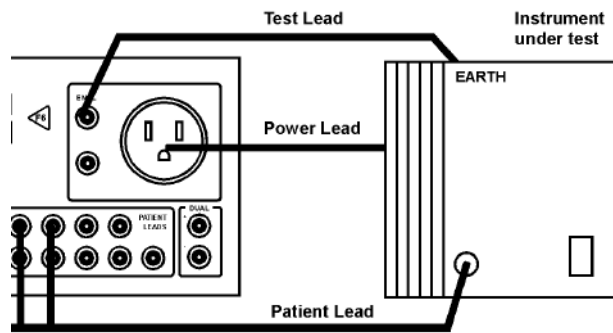


Figure 3 Equipment Leakage Current Test - Setup Example

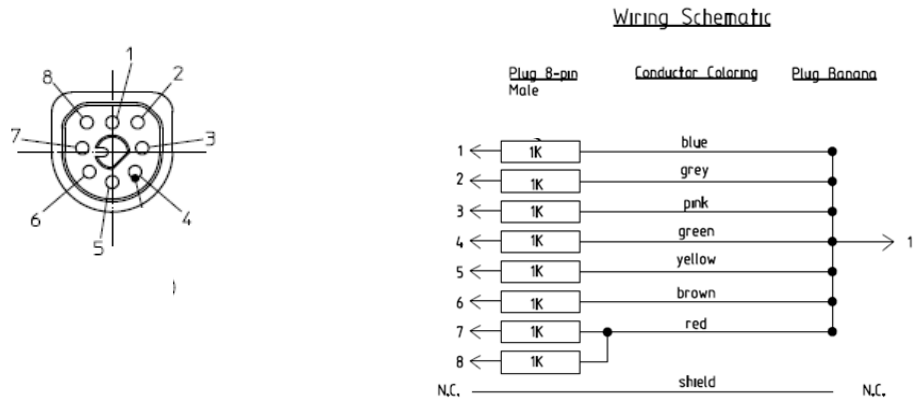


**Figure 4 Applied Part Current Test - Setup Example**

**NOTE** The above graphics resemble the Metron QA-90 setup and are protected by copyright. Copyright owned by Fluke (Metron).

### Safety Test Adapter Cable - Schematics

The following graphics provide schematics of safety test (patient lead) adapter cables which can be used for electrical safety testing. These schematics can also be used as a guideline for making your own safety test adapter cables. Alternatively, other methods to make safety test adapter cables can be used, e.g. using a modified accessory cable.



**Figure 5 FM20-FM50 Safety Test Adapter Cable Schematic**

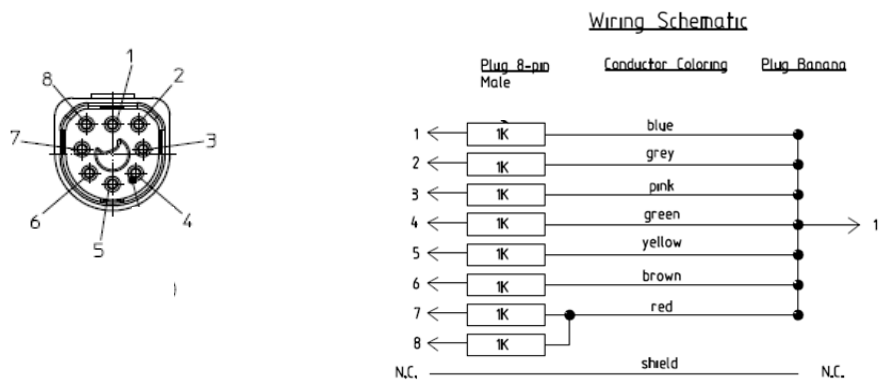


Figure 6 SpO<sub>2</sub> Safety Test Adapter Cable Schematic

### S(1): Detachable Power Cord Protective Earth Test (optional)

This test can be performed upon request by the customer.

Test to perform:

Use an Ohmmeter to measure the earth wire resistance of the detachable power cord.

This safety test is based on IEC/EN 60601-1, IEC/EN 62353, UL2601-1 Ed. 2/UL60601-1:2003 and CSA 601.1-M90.

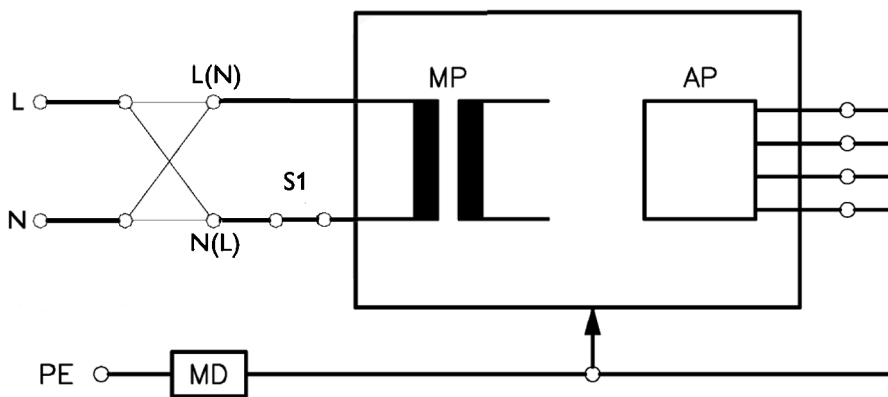
Report the highest value (X1).

Test	Expected Test Results
Protective Earth Resistance Test	X1 <= 100mOhms

- NOTE**
- If the protective earth resistance test fails, testing must be discontinued immediately and the device under test must be repaired or labeled as defective.
  - Flex the power cord during the protective earth resistance test to evaluate its integrity. If it does not pass the test, exchange the power cord.
  - The functional earth conductor is required for EMC purposes. It has no protective function against electrical shock. The protection against electrical shock is provided by double and/or reinforced insulation.

## S(2): Sum of Functional Earth and Equipment Leakage Current Test - Normal Condition

Test to perform:



**Measuring circuit for the measurement of Equipment Leakage Current - Direct method according to IEC/EN 62353.**

This test measures the leakage current of exposed metal parts of the FM20/FM30 monitor and the functional earth leakage current. It tests normal and reversed polarity. Perform the test with S1 closed (Normal Condition).

**NOTE** The protective earth conductor is required for EMC purposes. It has no protective function against electric shock. The protection against electric shock is provided by double and/or reinforced insulation.

*This safety test is based on IEC/EN 60601-1, IEC/EN 62353, and UL2601-1 Ed. 2/UL60601-1:2003 and CSA 601.1-M90.*

For measurement limits, refer to test block Safety (1), “Test and Inspection Matrix” on page 57.

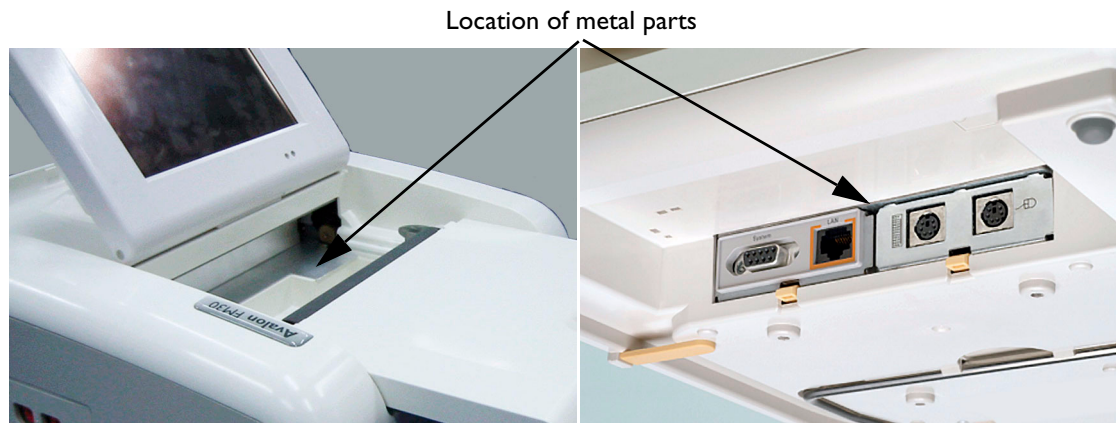
Report the highest value: (X1).

Test	Expected Test Results
Equipment Leakage Current Test (Normal Condition - with mains cable)	X1 <= 100µA

**NOTE** All values for current and voltage are the root mean square (r.m.s.) values, unless otherwise stated.

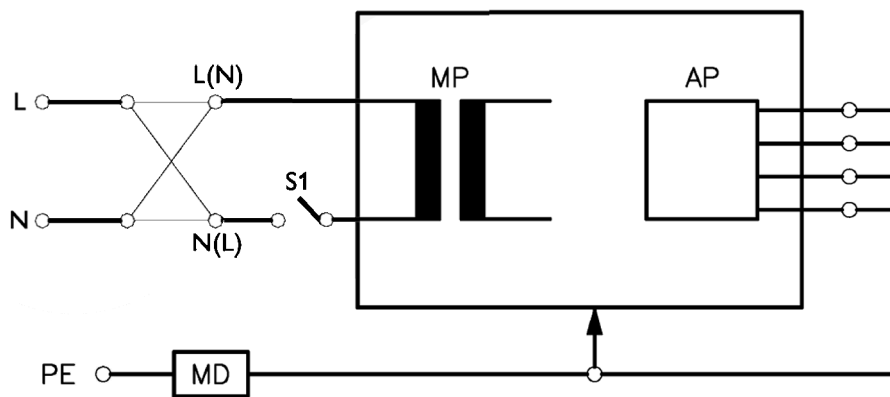
You can find metal parts of the device:

- In the top cover (accessed by opening the paper drawer).
- In an interface slot located in the bottom housing (you need to remove the interfaces if they are fitted).



### S(3): Sum of Functional Earth and Equipment Leakage Current Test - Single Fault Condition

Test to perform:



**Measuring circuit for the measurement of Equipment Leakage Current - Direct method according to IEC/EN 62353.**

This test measures leakage current of exposed metal parts of the FM20/FM30 monitor and the functional earth leakage current. It tests normal and reversed polarity. Perform the test with S1 open (Single Fault Condition).

**NOTE** The protective earth conductor is required for EMC purposes. It has no protective function against electric shock. The protection against electric shock is provided by double and/or reinforced insulation.  
*This safety test is based on IEC/EN 60601-1, IEC/EN 62353, and UL2601-1 Ed. 2/UL60601-1:2003 and CSA 601.1-M90.*

For measurement limits, refer to test block Safety (2), “Test and Inspection Matrix” on page 57.

Report the highest value: (X2).

Test	Expected Test Results
Equipment Leakage Current Test (Single Fault Condition - with mains cable)	X1 <= 300µA

**NOTE** All values for current and voltage are the root mean square (r.m.s.) values, unless otherwise stated.

You can find metal parts of the device:

- In the top cover (accessed by opening the paper drawer).
- In an interface slot located in the bottom housing (you need to remove the interfaces if they are fitted).

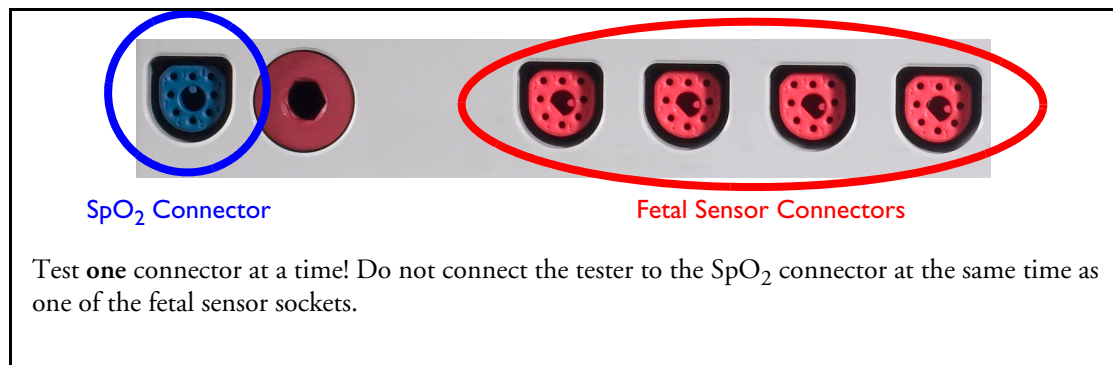
Refer to the picture showing accessible metal parts under “S(2): Sum of Functional Earth and Equipment Leakage Current Test - Normal Condition” on page 38.

## S(4): Applied Part Leakage Current, Mains on Applied Part

**NOTE** During measurement of the Applied Part Leakage Current it is possible that the measured current can exceed the allowed limit (per IEC/EN 60601-1 or IEC/EN 62353).

This can occur when the safety tester is connected to more than one connector simultaneously, that is, either to two fetal sensor connectors at the same time, or to a fetal sensor connector and the SpO<sub>2</sub> connector at the same time during the applied leakage current measurement.

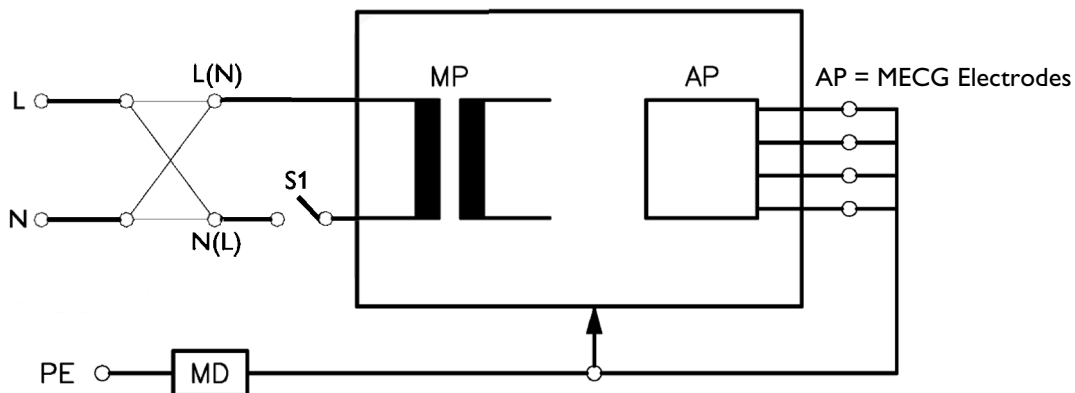
The connectors for the fetal sensors and for SpO<sub>2</sub> are independently functioning connectors.



Although there are individual connectors on the front end, internally those parameters use the same electrical insulation interface and are hardwired to each other. This results in an electrical short of those connectors during measurement if a test current is applied simultaneously. Therefore this should be avoided.

Due to the combined insulation interface, it is sufficient to connect to only one parameter interface (that is, one of the fetal sensor connectors or SpO<sub>2</sub>) of the fetal sensor/SpO<sub>2</sub> measurement block. This avoids a short and the potential of exceeding the limit for the current.

Test to perform:



Measuring circuit for the measurement of Applied Part Leakage Current - Direct method according to IEC/EN 62353.

This test measures applied part leakage current from applied part to earth caused by external main voltage on the applied part. Each polarity combination possible shall be tested. This test is applicable for ECG measurement inputs.

*This safety test is based on IEC/EN 60601-1, IEC/EN 62353, and UL2601-1 Ed. 2/UL60601-1:2003 and CSA 601.1-M90.*

For measurement limits and test voltage, refer to test block Safety (4), “Test and Inspection Matrix” on page 57.

Report the highest value: (X1).

Test	Expected Test Results
Applied Part Leakage Current Test (Single Fault Condition - mains on applied part)	X1 <= 50µA

**NOTE** All values for current and voltage are the root mean square (r.m.s.) values, unless otherwise stated.

## System Test

After mounting and setting up a system, perform system safety tests according to IEC/EN 60601-1-1.

### What is a Medical Electrical System?

A medical electrical system is a combination of at least one medical electrical device and other electrical equipment, interconnected by functional connection or use of a multiple portable socket-outlet.

- Devices forming a system must comply with IEC/EN 60601-1-1.
- Any electrical device such as IT equipment that is connected to the medical electrical equipment must comply with IEC/EN 60601-1-1 and be tested accordingly.

### General Requirements for a System

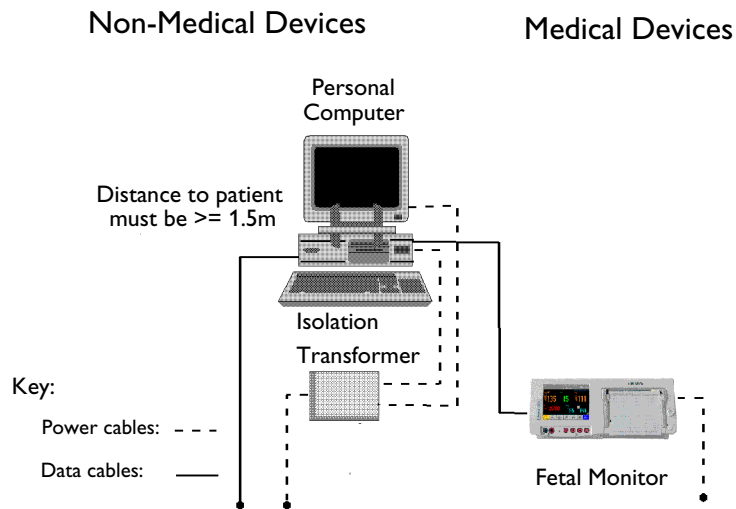
After installation or subsequent modification, a system must comply with the requirements of the system standard IEC/EN 60601-1-1. Compliance is checked by inspection, testing or analysis, as specified in the IEC/EN 60601-1-1 or in this book.

Medical electrical equipment must comply with the requirements of the general standard IEC/EN 60601-1, its relevant particular standards and specific national deviations. Non-medical electrical equipment shall comply with IEC and ISO safety standards that are relevant to that equipment.

Relevant standards for some non-medical electrical equipment may have limits for equipment leakage currents higher than required by the standard IEC/EN 60601-1-1. These higher limits are acceptable only outside the patient environment. It is essential to reduce equipment leakage currents when non-medical electrical equipment is to be used within the patient environment.

## System Example

This illustration shows a system where both the medical electrical equipment and the non-medical electrical equipment are situated at the patient's bedside.



- 
- WARNING**
- Do not use additional AC mains extension cords or multiple portable socket-outlets. If a multiple portable socket-outlet is used, the resulting system must be compliant with IEC/EN 60601-1-1. Do not place multiple socket-outlets on the floor. Do not exceed the maximum permitted load for multiple socket-outlets used with the system. Do not plug additional multiple socket-outlets or extension cords into multiple socket-outlets or extension cords used within the medical system
  - Do not connect any devices that are not supported as part of a system.
  - Do not use a device in the patient vicinity if it does not comply with IEC/EN 60601-1. The whole installation, including devices outside of the patient vicinity, must comply with IEC/EN 60601-1-1. Any non-medical device, **including a PC running an OB TraceVue system**, placed and operated in the patient's vicinity must be powered via a separating transformer (compliant with IEC/EN 60601-1-1) that ensures mechanical fixing of the power cords and covering of any unused power outlets.
- 

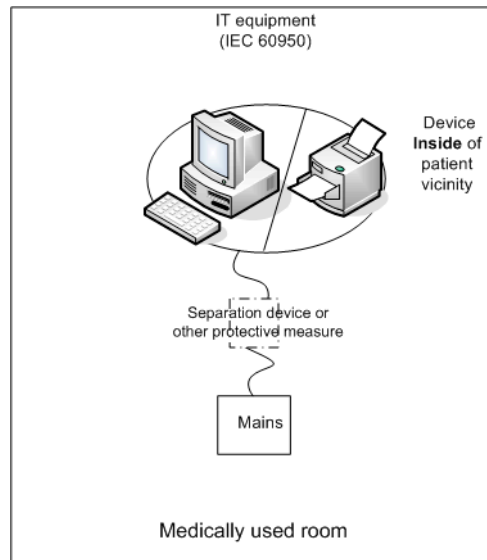
## System Installation Requirements

- Ensure that the medical electrical system is installed in a way that the user achieves optimal use.
- Make sure the user is informed about the required cleaning, adjustment, sterilization and disinfection procedures listed in the Instructions for Use.
- The medical electrical system must be installed in such a way that the user is able to carry out the necessary cleaning, adjustment, sterilization and disinfection procedures listed in the Instructions for Use.
- Ensure that the medical electrical system is installed in a way that an interruption and restoration of power to any part of the medical electrical system does not result in a safety hazard.

- We recommend using fixed mains socket outlets to power the medical system or parts thereof. Avoid using multiple portable socket-outlets.
- Any multiple portable socket outlets used must be compliant with IEC 60884-1 and IEC 60601-1-1.
- Ensure that any part of the system connected to multiple portable socket-outlets is only removable with a tool, i.e. the multiple portable socket-outlet provides a locking mechanism to prevent power cords from being plugged or unplugged unintentionally. Otherwise, the multiple portable socket-outlet must be connected to a separation device. Multiple Socket Outlets used within the medical electrical system must only be used for powering medical electrical equipment which is part of the system.
- Ensure that any functional connections between parts of the medical electrical system are isolated by a separation device according to IEC 60601-1-1 to limit increased equipment leakage currents caused by current flow through the signal connections. This only works if the equipment leakage current of the respective medical electrical system parts is not exceeded under normal conditions.
- Avoid increase of equipment leakage currents when non-medical electrical equipment within the medical electrical system is used. This only works if the equipment leakage current of the respective medical electrical system parts is not exceeded under normal conditions. Use additional protective earth connection, separation device or additional non-conductive enclosures.
- Within the patient environment it is important to limit electrical potential differences between different parts of a system. If necessary, use potential equalization equipment (equipotential cable) or additional protective earth connections.
- Medical electrical equipment used in medical rooms must be connected to potential equalization equipment (equipotential cable) to avoid electrical potential differences. Check your local requirements for details.

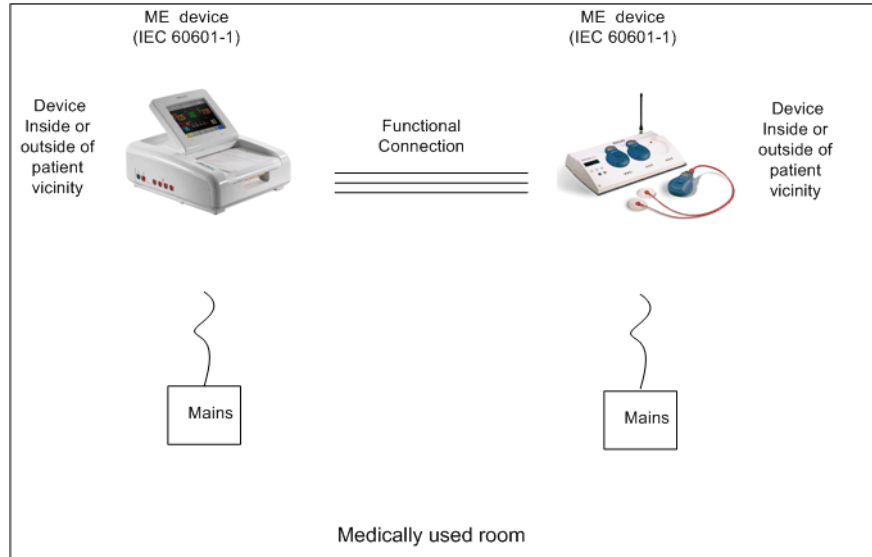
## Required Protective Measures at System Installation

For any IT equipment (IEC60950) operated in patient vicinity ensure that the equipment leakage current does not exceed the limits described in IEC 60601-1. Use a separation device to ensure compliance. After installation of IT equipment in patient vicinity, an enclosure leakage current test is required.

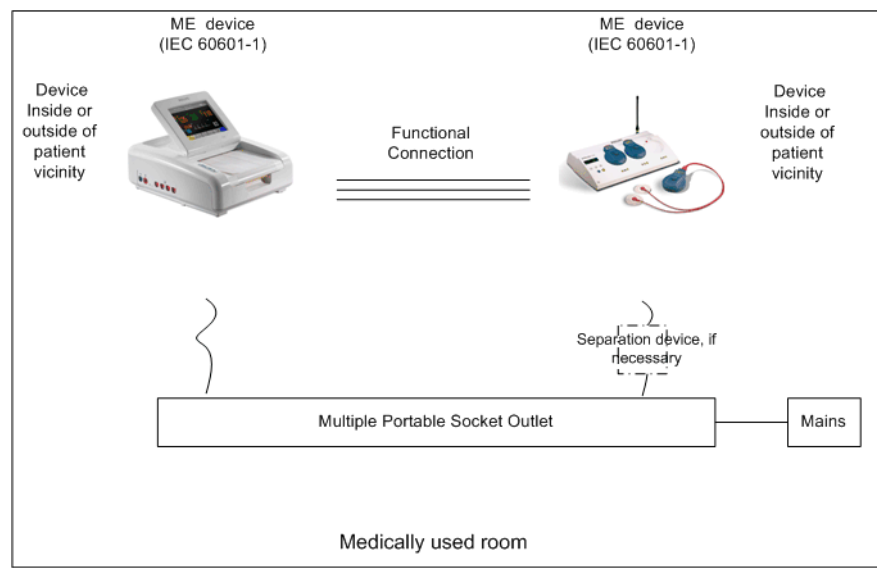


## Case 1: Medical Device Combined with Medical Device

If you combine a medical device with another medical device (incl. Philips specified displays) to form a medical electrical system according to IEC60601-1-1, no additional protective measures are required. The medical electrical devices may be located in or outside the patient vicinity in a medically used room. This is valid as long as the medical devices are connected to separate mains outlets. No system test is required.

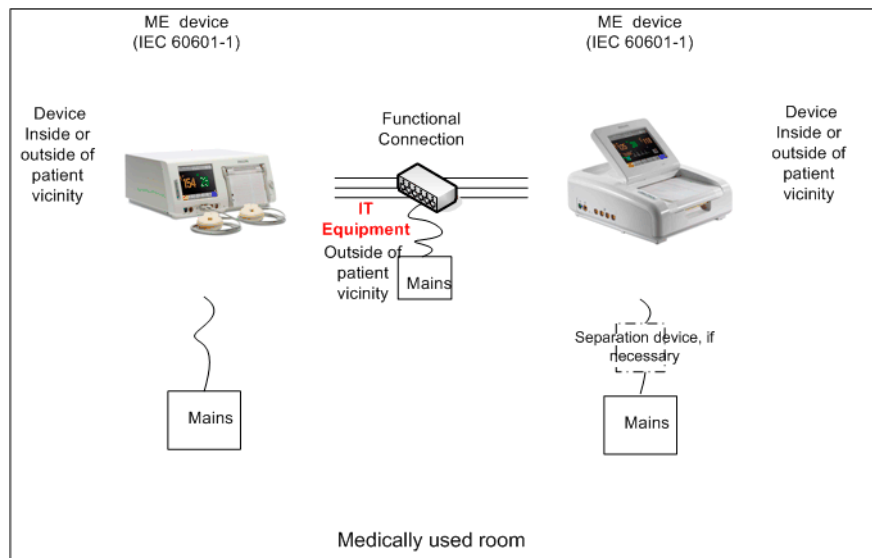


If the combined medical devices are connected to the same multiple portable socket outlet an enclosure leakage current test of the entire device combination on the multiple portable socket outlet is required to ensure that the resulting protective earth leakage current and equipment leakage current does not exceed the limits of IEC 60601-1-1. Avoid using multiple portable socket outlets. The medical electrical devices may be located in or outside the patient vicinity in a medically used room. If the limits are exceeded, additional protective measures are required, e.g. a separation device or the connection of each device to separate mains.

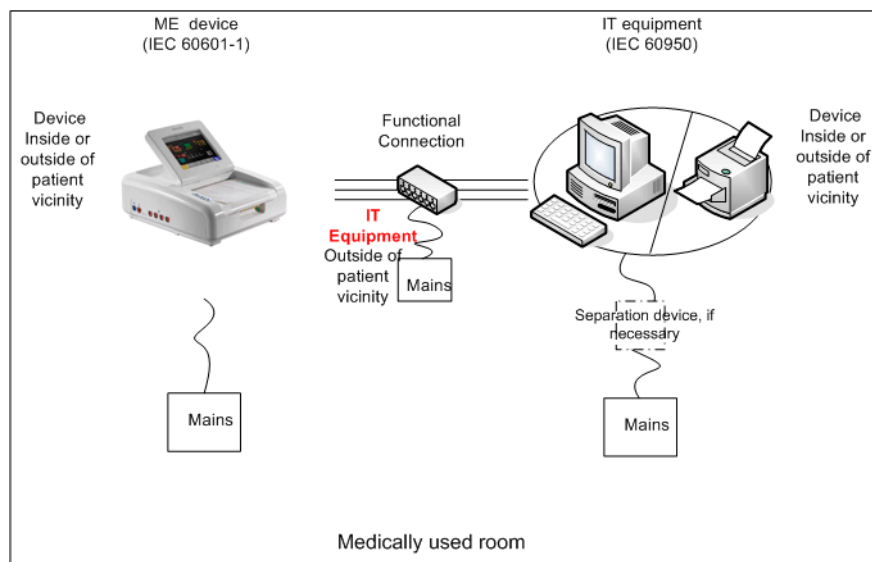


## Case 2: Medical Device Combined with a Non-Medical Device

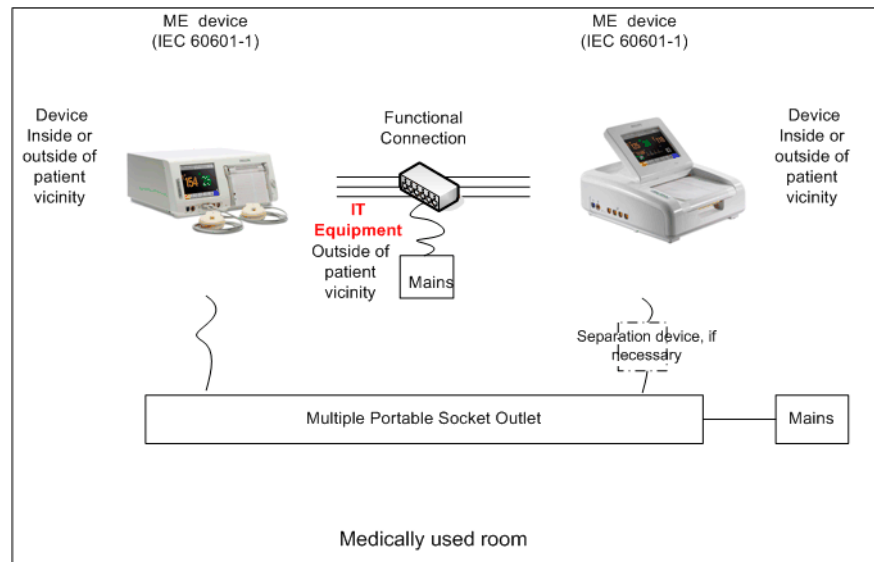
If you combine a medical device with a non-medical device to form a medical electrical system according to IEC60601-1-1, additional protective measures are required, e.g. usage of a separation device. The medical electrical devices or the IT equipment may be located in or outside the patient vicinity in a medically used room. After system installation incl. protective measures, a system test is required to ensure that the resulting equipment leakage current and applied part leakage current does not exceed the limits of IEC 60601-1-1.



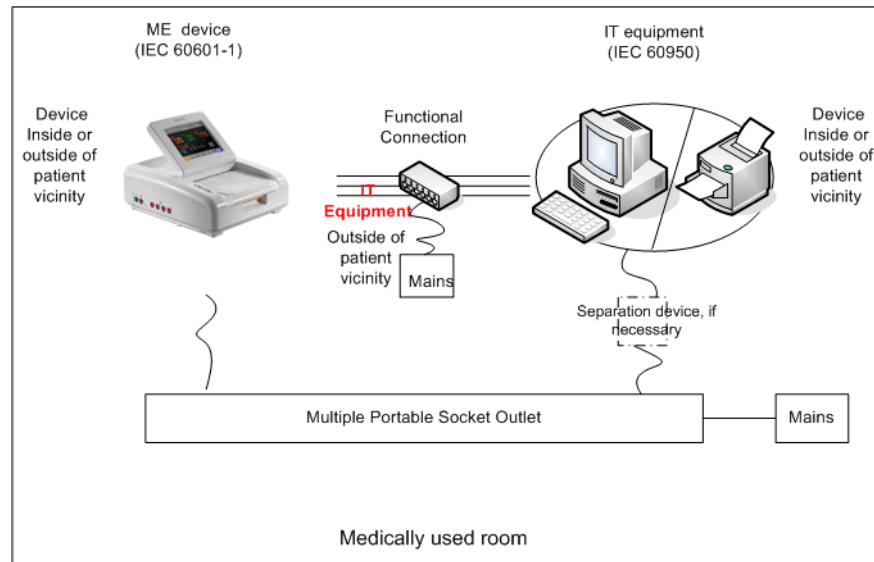
For any IT equipment (IEC60950) operated in patient vicinity ensure that the equipment leakage current does not exceed the limits described in IEC 60601-1. Use a separation device to ensure compliance. After installation of IT equipment in patient vicinity, an enclosure leakage current test is required.



If the combined devices forming the medical electrical system are connected to the same multiple portable socket outlet, ensure that the resulting protective earth leakage current **and** equipment leakage current do not exceed the limits of IEC 60601-1-1. The medical electrical devices or IT equipment may be located in or outside the patient vicinity in a medically used room. Avoid using multiple portable socket outlets. If the limits of IEC 60601-1-1 are exceeded, additional protective measures are required, e.g. a separation device or the connection of each device to separate mains.

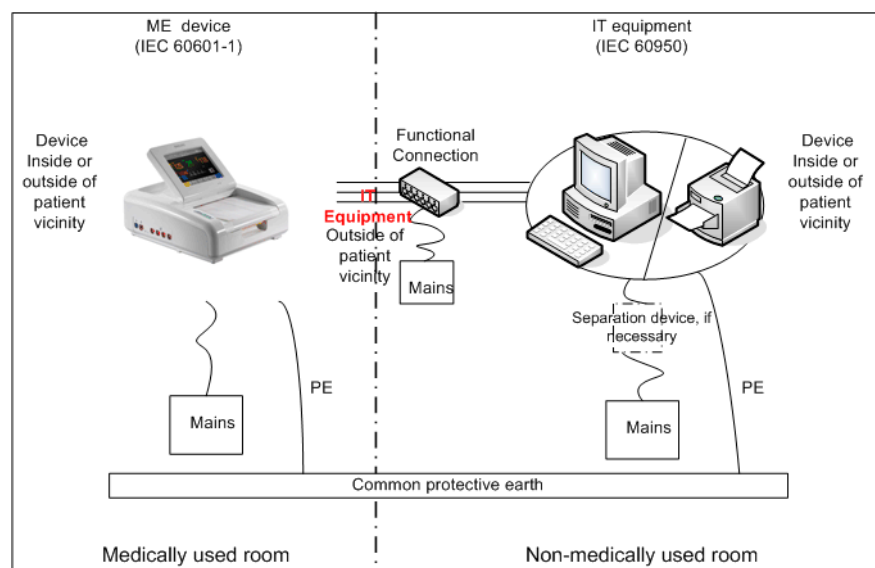
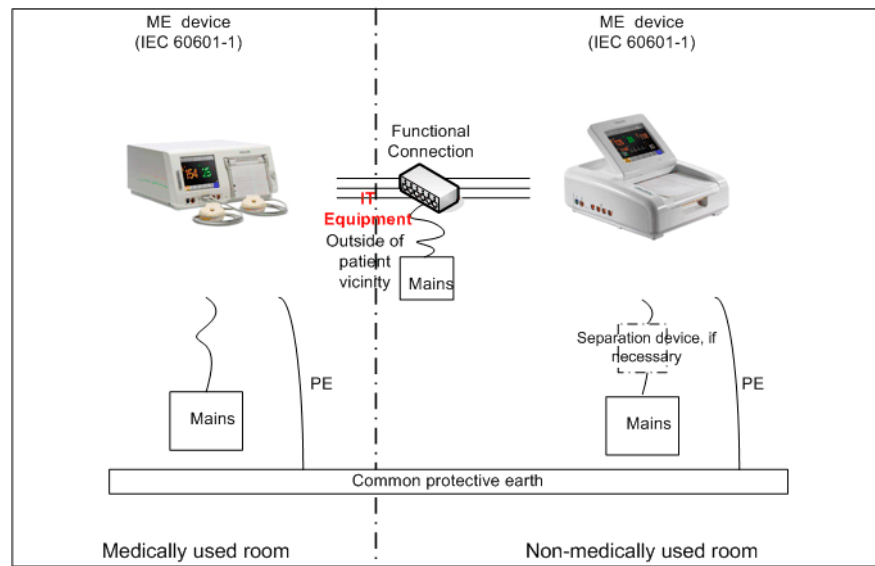


For any IT equipment (IEC 60950) operated in patient vicinity, ensure that the equipment leakage current does not exceed the limits described in IEC 60601-1. Use a separation device to ensure compliance. After installation of IT equipment in patient vicinity, an enclosure leakage current test is required.

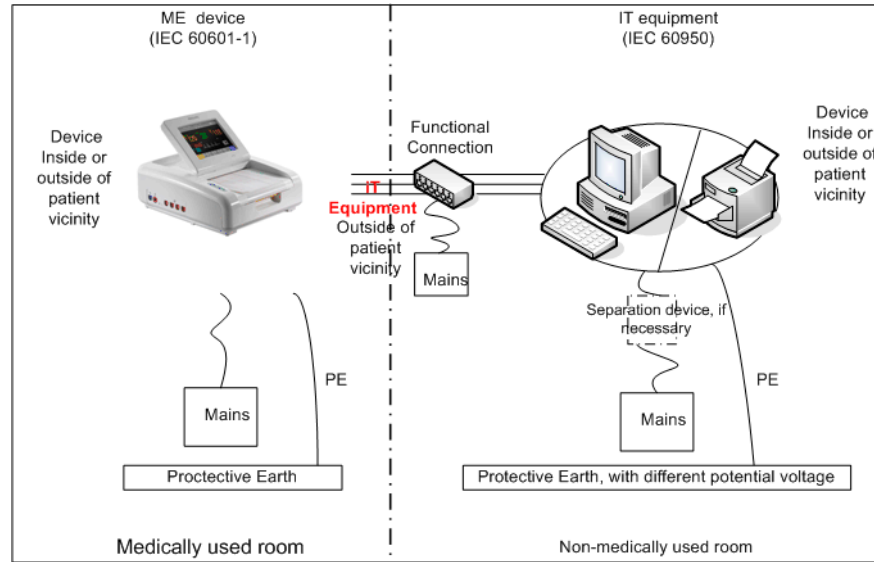
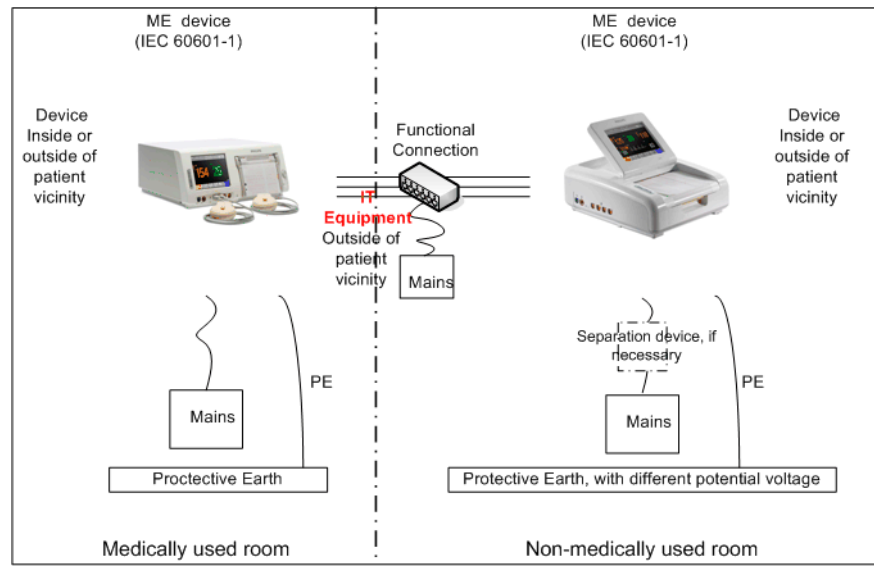


### Case 3: Medical Device Combined with a Medical or Non-Medical Device with one Device in a Non-Medically-Used Room

If you combine a medical device with a medical or non-medical device to form a medical electrical system according to IEC60601-1-1 using a common protective earth connection and one of the devices is located in a non-medically used room, additional protective measures are required, e.g. usage of a separation device or additional protective earth connection. The medical electrical devices or IT equipment may be located in or outside the patient vicinity. After system installation incl. protective measures, a system test is required to ensure that the resulting equipment leakage current does not exceed the limits of IEC 60601-1-1.



If you combine a medical device with a medical or non-medical device to form a medical electrical system according to IEC60601-1-1 using two separate protective earth connections and one of the devices is located in a non-medically used room creating a potential voltage difference, additional protective measures are required, e.g. usage of a separation device or additional protective earth connection. The medical electrical devices or IT equipment may be located in or outside the patient vicinity. After system installation incl. protective measures, a system test is required to ensure that the resulting equipment leakage current does not exceed the limits of IEC 60601-1-1.



## System Test Procedure

If the medical electrical device has already been tested as a standalone device e.g. during factory safety testing, an equipment leakage current test must be performed once the device is connected to the LAN network. If the medical electrical device has not been tested as a standalone device, the device has to be tested as a standalone device (without connection to the system) and as part of the system (with connection to the system).

Connect the detachable power cord of the device under test to the safety analyzer's test mains port. Connect the enclosure test lead of the safety analyzer to the enclosure of the device under test, e.g. to the equipotential connector. Refer to the documentation that accompanies the safety analyzer for further details on how to set up the test.

Test	Expected Test Results
Equipment Leakage Current Test (Normal Condition)	Sys1 <= 100µA
Equipment Leakage Current Test (Single Fault Condition)	Sys2 <= 300µA

After the testing of the device as a standalone device and as part of the system, check that the resulting values (without connection and with connection to the system) do not differ by more than +/- 10% from each other.

If the devices in the medical electrical system are connected to a multiple portable socket outlet the resulting protective earth leakage current needs to be determined. All system components must be connected to the multiple portable socket outlet and be switched on during this measurement.

Test	Expected Test Results
Protective Earth Leakage Current of Multiple Socket-Outlets	Sys3 <= 100µA

Refer to the documentation that accompanies the safety analyzer for further details on how to set up the test.

## Preventive Maintenance Procedures

The preventive maintenance tasks are restricted to the noninvasive blood pressure measurement calibration. Carry out the noninvasive blood pressure measurement performance tests at least every two years, or as specified by local laws (whichever comes first).

### Noninvasive Blood Pressure Measurement Calibration

Carry out the noninvasive blood pressure measurement performance tests at least every two years, or as specified by local laws (whichever comes first).

## Fetal Recorder Maintenance

The recorder rubber roller, thermal printhead and paper sensor should be cleaned at least once a year, or when needed (when traces become faint).

Clean the assemblies as follows:

- Clean the recorder rubber roller with a lint-free cloth using a soap/water solution.
- Wipe the printhead using a cotton swab moistened with 70% Isopropyl alcohol based solution.
- Check the paper sensing mechanism is dust free.

## Performance Assurance Tests

Some of the following test procedures must be performed in service mode. To enter service mode select **Operating Modes** in the main menu. Then select **Service Mode** and enter the password.

### Basic Performance Assurance Test

This section describes the basic performance test procedure. Please refer to the section *When to Perform Tests* for detailed information on when which test procedure is required.

**Procedure:**

Power on the monitoring system and go into demo mode. Check that each parameter displays values.

### Full Performance Assurance Test

The following sections describe the full performance testing procedures i.e. detailed testing of each parameter with a patient simulator or specified tools. Please refer to the section *When to perform Tests* for information on when which testing procedure is required.

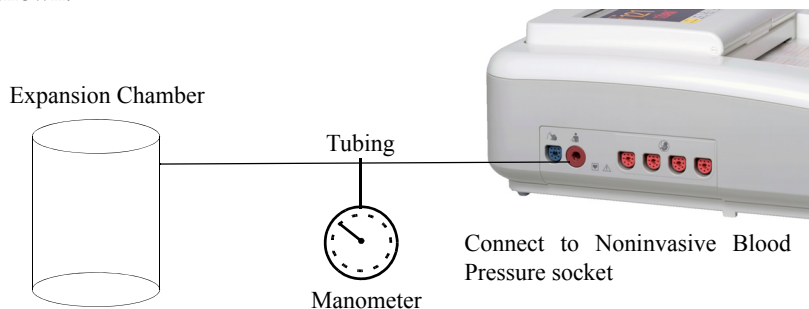
### Noninvasive Blood Pressure Performance Tests

This section describes noninvasive blood pressure test procedures. The monitor must be in service mode.

Table 3 gives the expected test results for each of the tests.

#### Accuracy Test

This test checks the performance of the noninvasive blood pressure measurement. Connect the equipment as shown:



Tools required:

- Reference manometer (includes hand pump and valve), accuracy 0.2% of reading.
- Expansion chamber (volume 250 ml +/- 10%)
- Appropriate tubing.

In service mode, the systolic and diastolic readings indicate the noise of noninvasive blood pressure channels 1 and 2 respectively. When static pressure is applied, the reading in noninvasive blood pressure channel 1 should be below 50. The value in parentheses indicates the actual pressure applied to the system.

- 1 Connect the manometer and the pump with tubing to the noninvasive blood pressure connector on the monitor and to the expansion chamber.
- 2 In service mode, select the **Setup NBP** menu.
- 3 Select **Close Valves: On**
- 4 Raise the pressure to 280 mmHg with the manometer pump.
- 5 Wait 10 seconds for the measurement to stabilize.
- 6 Compare the manometer values with the displayed values.
- 7 Document the value displayed by the monitor (X1).
- 8 If the difference between the manometer and displayed values is greater than 3 mmHg, calibrate the noninvasive blood pressure measurement. If not, proceed to the leakage test.
- 9 To calibrate the noninvasive blood pressure measurement, select **Close Valves off** then **Calibrate NBP** and wait for the instrument to pump up the expansion chamber. Wait a few seconds after pumping stops until **EnterPrVal** is highlighted and then move the cursor to the value shown on the manometer. If one of the following prompt messages appears during this step, check whether there is leakage in the setup:
  - NBP unable to calibrate—cannot adjust pressure
  - NBP unable to calibrate—unstable signal
- 10 Press **Confirm**.

If the INOP NBP Equipment Malfunction message occurs in monitoring mode, go back to service mode and repeat the calibration procedure.

## Leakage Test

The noninvasive blood pressure leakage test checks the integrity of the system and of the valve. It is required once every two years and when you repair the monitor or replace parts.

- 1 If you have calibrated, repeat steps 2 to 6 from the accuracy test procedure so that you have 280 mmHg pressure on the expansion chamber.
- 2 Watch the pressure value for 60 seconds.
- 3 Calculate and document the leakage test value (X2).  
$$X2 = P1 - P2$$
where P1 is the pressure at the beginning of the leakage test and P2 is the pressure displayed after 60 seconds.  
The leakage test value should be less than 6 mmHg.

## Linearity Test

- 1 Reduce the manometer pressure to 150 mmHg.
- 2 Wait 10 seconds for the measurement to stabilize.
- 3 After these 10 seconds, compare the manometer value with the displayed value.
- 4 Document the value displayed by the monitor (X3)
- 5 If the difference is greater than 3 mmHg, calibrate the noninvasive blood pressure measurement (see steps 9 to 10 in the accuracy test procedure).

## Valve Test

- 1 Raise the pressure again to 280 mmHg.
- 2 Select **Close valves: Off**.
- 3 Wait five seconds and then document the value displayed. The value should be less than 10 mmHg.
- 4 Document the value displayed by the monitor (X4).

## Expected Test Results

Test	Expected test results
Accuracy test	x1 = value displayed by monitor Difference ≤ 3mmHg
Leakage test	x2 = leakage test value x2 < 6 mmHg
Linearity test	x3 = value displayed by monitor Difference ≤ 3mmHg
Valve Test	x4 = value < 10 mmHg

## SpO<sub>2</sub> Performance Test

This test checks the performance of the SpO<sub>2</sub> measurement.

**Tools required:** none

- 1 Connect an adult SpO<sub>2</sub> transducer to the SpO<sub>2</sub> connector.
- 2 Measure the SpO<sub>2</sub> value on your finger (this assumes that you are healthy).
- 3 The value should be between 95% and 100%.

## Expected Test Results

Test	Expected test results
SpO <sub>2</sub> Performance Test	95% and 100%

## Measurement Validation

The SpO<sub>2</sub> accuracy has been validated in human studies against arterial blood sample reference measured with a CO-oximeter. In a controlled desaturation study, healthy adult volunteers with saturation levels between 70% and 100% SaO<sub>2</sub> were studied.

The population characteristics for those studies were:

- Gender: about half female and half male subjects
- Age range: 18 to 45 years
- Skin tone range: from light to very dark

**NOTE** A functional tester cannot be used to assess the accuracy of a pulse oximeter monitor. However, it can be used to demonstrate that a particular pulse oximeter monitor reproduces a calibration curve that has been independently demonstrated to fulfill a particular accuracy specification.

## Reporting of Test Results

Philips recommends all test results are documented in accordance with local laws. Authorized Philips personnel report test result back to Philips to add to the product development database. While hospital personnel (biomedical engineers or technicians) do not need to report results to Philips, Philips recommends that they record and store the test results in accordance with local laws.

The following table lists what to record after completing the tests in this chapter. Record the results in the empty column in the following table.

The following is a guide as to what your documentation should include:

- Identification of the testing body (for example, which company or department carried out the tests).
- Name of the person(s) who performed the tests and the concluding evaluation.
- Identification of the device(s) and accessories being tested (serial number, etc.).
- The actual tests (incl. visual inspections, performance tests, safety and system tests) and measurements required
- Date of testing and of the concluding evaluation.
- A record of the actual values of the test results, and whether these values passed or failed the tests.
- Date and confirmation of the person who performed the tests and evaluation.

The device under test should be marked according to the test result: passed or failed.

## Carrying Out and Reporting Tests

### Test Report

Testing Organization:	Test before putting into service (reference value)	O
Name of testing person:	Recurrent Test	O
	Test after Repair	O
Responsible Organization:		
Device Under Test:	ID Number:	
Product Number	Serial No.:	
Accessories:		
Measurement Equipment (Manufacturer, Type, Serial No.):		
Functional Test (parameters tested):		

### Test and Inspection Matrix

**Table 3: Test and Inspection Matrix**

Test Block	Test or Inspection to be Performed	Expected Test Results	Record the Results (mandatory for Philips Personnel only)	
			What to Record	Actual Results
Visual Inspection	Perform Visual Inspection (see "Visual Inspection" on page 31).	Pass or Fail	V:P or V:F	
Power On	Power on the unit. Does the self-test complete successfully?	If Yes, Power On test is passed.	PO:P or PO:F	
Noninvasive Blood Pressure Performance Tests	Perform the Accuracy Test (see page 53).	X1 = value displayed by monitor Difference <= 3mmHg	PN:P/X1 or PN:F/X1	
	Performance Leakage Test (see page 54).	X2 = leakage test value X2 < 6 mmHg	PN:P/X2 or PN:F/X2	
	Performance Linearity Test (see page 55).	X3 = value displayed by monitor Difference <= 3mmHg	PN:P/X3 or PN:F/X3	
	Performance Valve Test (see page 55).	X4 = value < 10 mmHg	PN:P/X4 or PN:F/X4	
SpO <sub>2</sub> Performance Test	Perform the SpO <sub>2</sub> Performance Test (see page 55).	Value should be between 95% and 100%	No reporting necessary	
Safety (2)	Perform Safety Test (2): Sum of Functional Earth and Equipment Leakage Current - Normal Condition.	NC with mains cable: Maximum leakage current (X1) ≤ 100µA	S(2):P/X1 or S(2):F/X1	

**Table 3: Test and Inspection Matrix**

Test Block	Test or Inspection to be Performed	Expected Test Results	Record the Results (mandatory for Philips Personnel only)	
			What to Record	Actual Results
Safety (3)	Perform Safety Test (3): Sum of Functional Earth and Equipment Leakage Current - Single Fault Condition (Open Earth).	SFC with mains cable: Maximum leakage current (X2) $\leq 300\mu\text{A}$	S(3):P/X2 or S(3):F/X2	
Safety (4)	Perform Safety Test (4): Applied Part Leakage Current - Single Fault Condition, mains on applied part.	Maximum leakage current (X1) $\leq 50\mu\text{A}$	S(4):P/X1 or S(4):F/X1	
System (Sys 1-2)	Perform the system test according to sub clause 19.201 of IEC/EN 60601-1-1, if applicable, after forming a system.	Equipment Leakage Current: Sys1 $\leq 100\mu\text{A}$ (Normal Condition) Sys2 $\leq 300\mu\text{A}$ (Single Fault Condition)	Sys: PSys1/PSys2 or Sys: FSys1/FSys2	
System (Sys3)	Perform the system test according to sub clause 19.201 of IEC/EN 60601-1-1, if applicable, after forming a system.	Protective Earth Leakage Current if medical electrical system components are connected to the same Multiple Portable Socket-Outlet: Sys3 $\leq 300\mu\text{A}$	Sys: PSys3 or Sys: FSys3	

Key: P = Pass, F = Fail, X or Sys = test result value to be recorded

**NOTE** All values for current and voltage are the root mean square (r.m.s.) values, unless otherwise stated.

## Evaluation

	Yes	No
Safety and Functional Test passed		
Repair required at a later date, safety and functional test passed		
Device must be taken out of operation until repair and passed tests		
Device failed and must be taken out of operation		
Notes:		
Next Recurrent Test:		
Name:		
Date/Signature:		

## Evaluation of Test Results

The evaluation of the test results must be performed by appropriately trained personnel with sufficient product, safety testing and application knowledge.

If any test results are between 90% and 100% of the respective expected result, the previously measured reference values must be taken into consideration for the assessment of the electrical safety of the device under test. If no reference values are available, you should consider shorter intervals between upcoming recurrent tests.

**NOTE** If any single test fails, testing must be discontinued immediately and the device under test must be repaired or labeled as defective. Be sure to inform the user about the test failure in writing.

## Other Regular Tests

The care and cleaning requirements that apply to the monitor and its accessories are described in the *Instructions for Use*. This section details the periodic maintenance recommended for the monitor, transducers and accessories.

## Transducers and Patient Modules; Functional Tests

If any of the following tests fail, repeat the test using another transducer. If the second transducer passes the tests, confirming that the first transducer is defective, contact your service personnel.

If the second transducer also fails the tests, contact your Philips Service Engineer or Response Center.

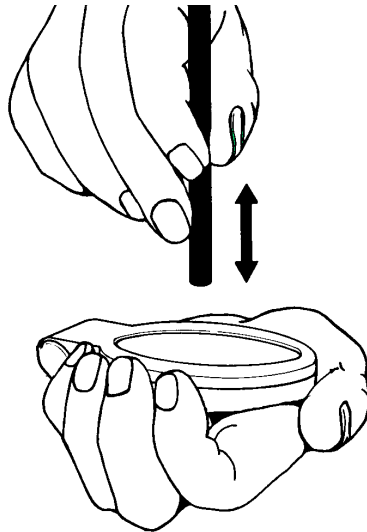
### Ultrasound Transducer Electrical Check

**CAUTION** Use of ultrasound gel that is not approved by Philips may reduce signal quality and may damage the transducer. This type of damage is not covered by warranty.

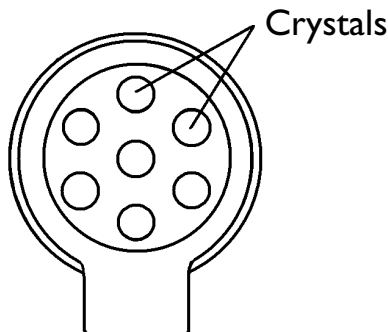
To test an ultrasound transducer:

- 1 Switch on the monitor and the recorder.
- 2 Connect the transducer to the fetal monitor.
- 3 Select the fetal heart sound for this channel.
- 4 Increase the loudspeaker volume to an audible level.

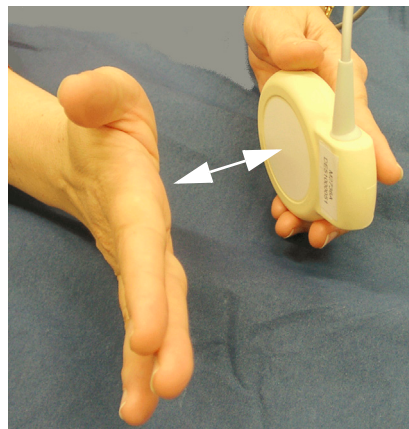
- 5 The ultrasound transducer contains seven piezoelectric crystals. Basic functioning of each can be verified by holding a flat bottomed pencil or similar above each crystal and moving it up and down as shown.



- 6 A sound should be heard for each crystal tested. The pencil should be held two to three centimeters from the transducer surface when the test is carried out.



- 7 A sound should also be heard when the transducer is moved back and forth over a solid surface, or the hand as shown.



## Toco Transducer Electrical Check

To test a Toco transducer:

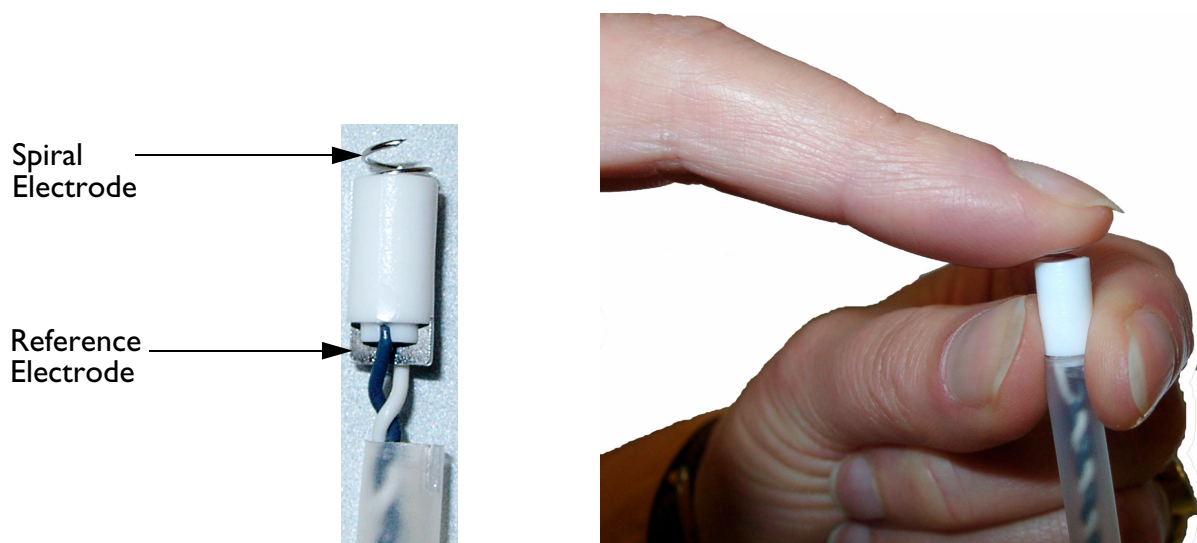
- 1 Switch on the monitor and the recorder.
- 2 Connect the transducer to the fetal monitor.
- 3 Lay the transducer face up on a hard, flat surface for a few seconds.
- 4 Press the Toco Baseline Key to re-adjust the Toco display to  $20\pm1$ .
- 5 Turn the transducer over so that the Toco sensor is resting on the flat surface. You should see a marked increase in the value of the Toco reading in the Toco display.
- 6 Press the Toco Baseline Key to re-adjust the Toco display to  $20\pm1$ .
- 7 Turn the transducer over again. You should see a marked decrease in the value of the Toco reading in the Toco display



## Testing the Patient Module (M2738A)/Toco<sup>+</sup> Transducer (M2735A): DECG Mode

- 1 Switch on the monitor and the recorder.
- 2 Connect the patient module or Toco<sup>+</sup> transducer to the fetal monitor.
- 3 Attach the DECG adapter cable M1362B to the socket on the patient module or Toco<sup>+</sup> transducer.
- 4 Ensure that the DFHR channel display on the fetal monitor shows the **DECG LEADS OFF** INOP with the DECG adapter cable attached.
- 5 Take a Fetal Scalp Electrode, and connect it to the DECG adapter cable.
- 6 *EITHER*  
Hold the reference electrode between the thumb and index finger of one hand, and touch the spiral electrode with the index finger of the other hand, as illustrated below. This makes a short between the spiral electrode and the reference electrode (it is best to wet your fingers first). Use a **sterile** Fetal Scalp Electrode.

**CAUTION** The tip of the spiral electrode is sharp. Take care not to injure your fingers.



*OR*

Cut off the plastic tip of the fetal scalp electrode (containing the spiral and reference electrodes) from the end of the wires. Strip the insulation from the end of the wires, and connect them to a patient simulator.

*Note*—We do not recommend the use of a specific patient simulator. The use of a patient simulator does not allow checking the specification of the ECG-functionality; it allows only a check of general function.

**Result:** the **DECG LEADS OFF INOP** should disappear.

**Viewing the ECG wave:** when configured, you can view the DECG wave on the screen, and any noise will be visible as additional verification of the effectiveness of the test.

If the test results are not as outlined above, repeat the test with another ECG transducer. If this does not solve the problem, try the following:

- Check all connections.
- If the **DECG LEADS OFF INOP** is still displayed, the DECG adapter cable may be defective. Replace the adapter cable.

If the problem persists, replace the transducer.

### Testing the Patient Module (M2738A)/Toco<sup>+</sup> Transducer (M2735A): MECG Mode

- 1 Switch on the monitor and the recorder.
- 2 Connect the patient module or Toco<sup>+</sup> transducer to the fetal monitor.
- 3 Attach the MECG adapter cable M1363A to the red color-coded socket on the patient module or Toco<sup>+</sup> transducer

**4 EITHER**

Attach electrodes to the M1363A adapter cable, and apply the electrodes to the skin (for example on the wrists).

*OR*

Attach the M1363A adapter cable to a patient simulator.

*Note*—We do not recommend the use of a specific patient simulator. The use of a patient simulator does not allow checking the specification of the ECG-Functionality; it allows only a check of general function.

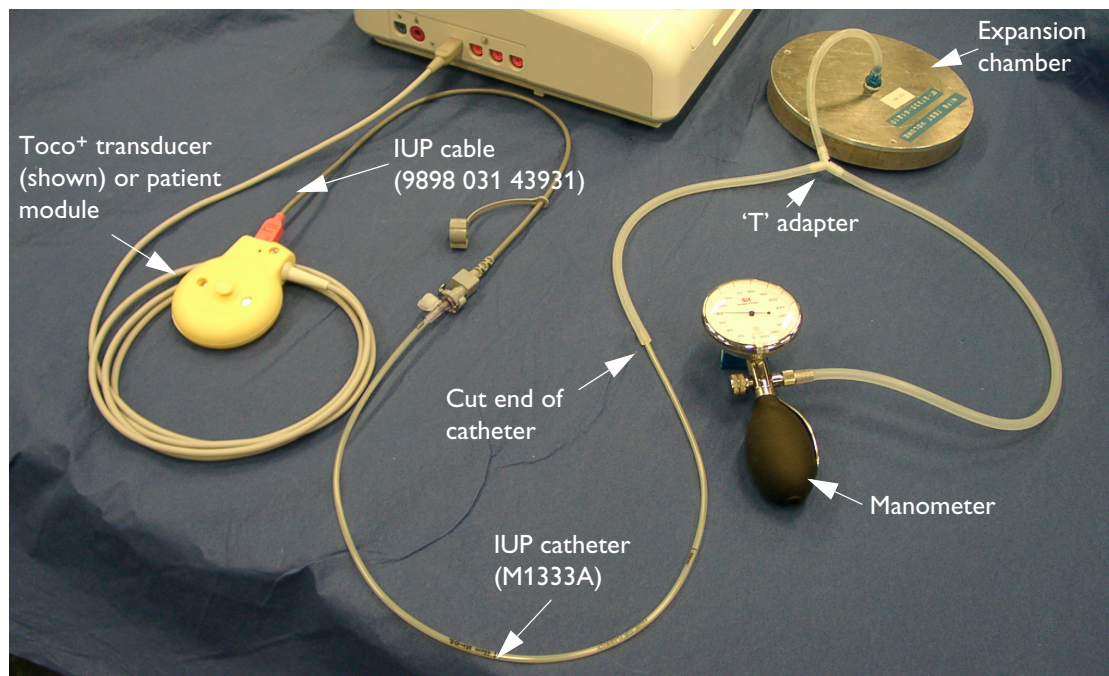
**Result:** You should see MEKG values displayed on the maternal display or annotated on the recorder trace.

If the test results are not as outlined above, repeat the test with another ECG transducer. If this does not solve the problem:

- The MEKG adapter cable may be defective. Replace the adapter cable, and repeat the test.
- Check all connections.

### Testing the Patient Module (M2738A)/Toco<sup>+</sup> Transducer (M2735A): IUP Mode

To test the IUP functionality of the patient module or the Toco<sup>+</sup> transducer, you need the following:



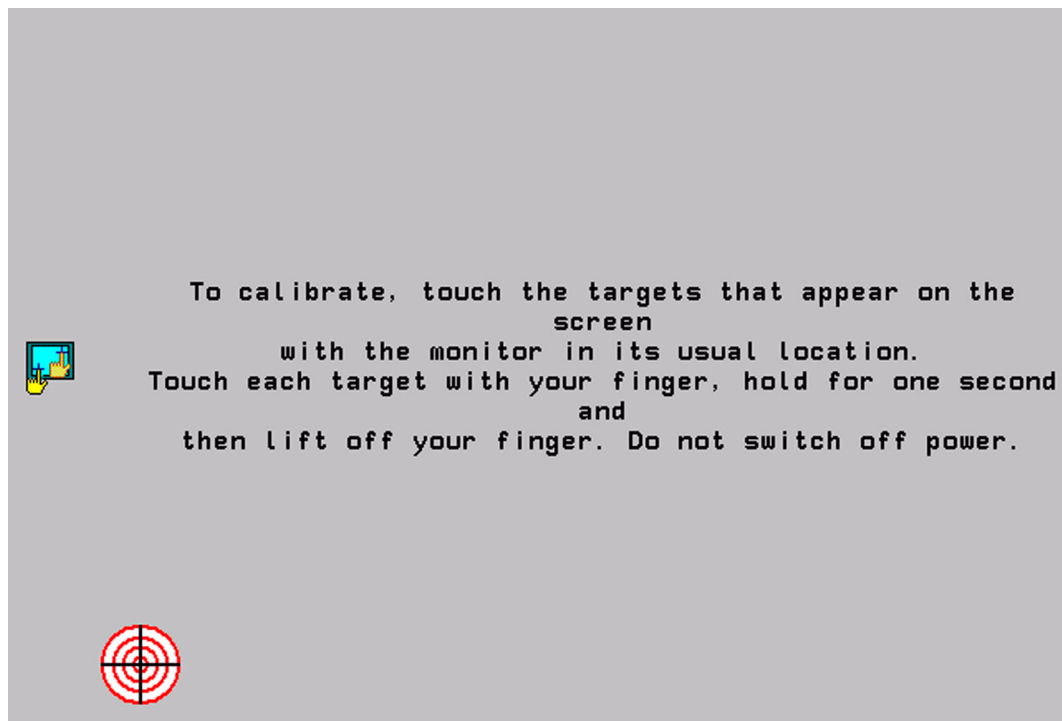
- Manometer.
  - Expansion chamber.
  - Three lengths of silicone tubing with a 'T' adapter.
- 1 Switch on the monitor and the recorder.
  - 2 Connect the patient module or Toco<sup>+</sup> transducer to the fetal monitor.
  - 3 Attach the IUP adapter cable ((989803143931)) to the socket on the patient module or Toco<sup>+</sup> transducer.

- 4 Cut the sensor tip off an IUP catheter (M1333A).
- 5 Connect the catheter to the IUP adapter cable.
- 6 Connect the silicone tubing to the test volume chamber and the manometer as shown in the picture.
- 7 Connect the cut end of the catheter to the silicone tubing.
- 8 Apply a pressure of  $80 \text{ mmHg} \pm 5 \text{ mmHg}$  with the manometer. Check that the value on the display and on trace corresponds to this pressure. Slowly release the pressure, and check that the value on the display and on trace shows this change in pressure.

## Touchscreen Calibration

To access the touchscreen calibration screen:

- 1 Enter service mode
- 2 Select **Main Setup**
- 3 Select **Hardware**
- 4 Select **Calibrate Touch**



Make sure you complete the calibration procedure without powering off the monitor mid-way. If the monitor is powered off after the first point is touched, the touch panel will be deactivated until the touch calibration is performed again.

If the touchscreen is accidentally mis-calibrated by selecting the wrong spot, you must use another input device to re-enter calibration mode, for instance, a mouse connected to the PS/2 interface. If you have the support tool, you can select **Start Touch Calibration** and it will create a rough calibration which will allow you to access the calibration menu again via the touchscreen.

## Disabling/Enabling Touch Operation

To disable touchscreen operation of the monitor, press and hold the **Main Screen** key for about three seconds. A red padlock will blink on the key. Press and hold the **Main Screen** key again for about three seconds to re-enable touchscreen operation.

## Checking the Fetal Recorder Offset

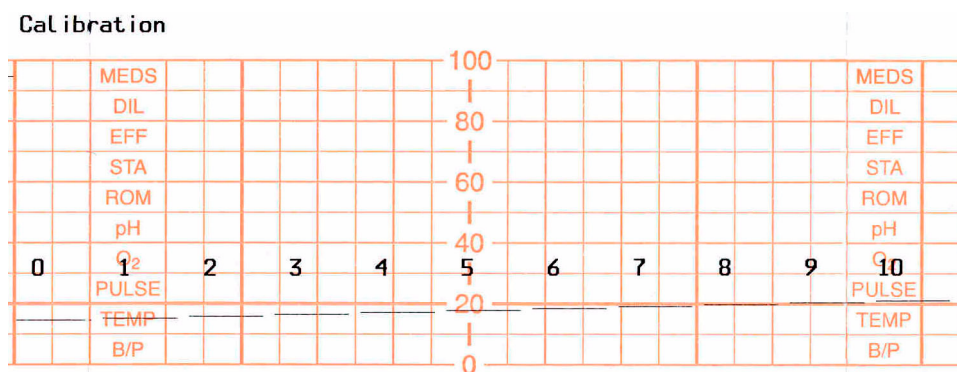
To check the recorder offset:

- 1 Connect a Toco transducer to the monitor.
- 2 Place the Toco transducer on a flat surface so that it is resting on the belt button, and is therefore not under any load.
- 3 Start the recorder, and press the Paper Advance key three times to make sure that at least three pages of paper have advanced.
- 4 If the Toco trace is recording exactly on the 20 unit gridline, then the offset is correctly set.
- 5 If the Toco trace is not recording exactly on the 20 unit gridline, then set the offset as described in “Setting the Fetal Recorder Offset”.

## Setting the Fetal Recorder Offset

To set the fetal recorder offset, you first need to run the fetal recorder calibration:

- 1 Enter Service Mode.
- 2 In **Main Setup**, select **Fetal Recorder** to enter the Fetal Recorder menu.  
The current setting for the recorder offset is shown (but it is still grayed out, and you cannot select it yet).
- 3 Select **Calibration** to start the recorder calibration printout.
- 4 The recorder stops, and the **Cal. Offset** becomes selectable.
- 5 Look at the section of the printout entitled “Calibration”.



You will see the numbers 0 to 10, and each number has a line printed below it. See for which number the line best fits the 20 unit gridline (8 in this example).

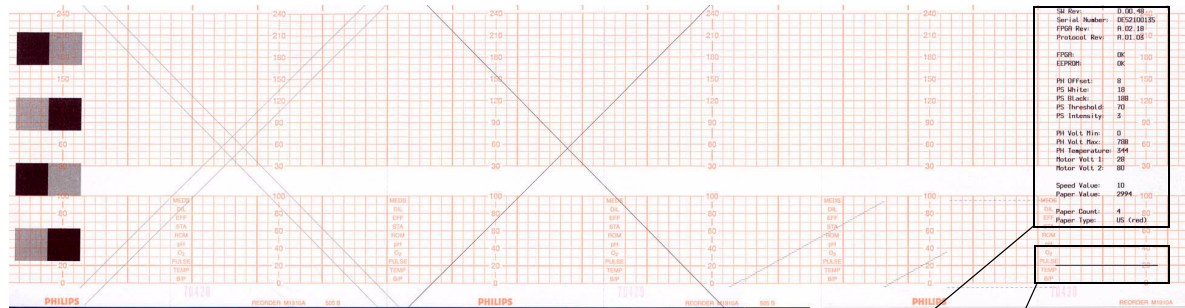
- 6 Select **Cal. Offset**, and select the offset value from 0 to 10 from the list, as determined in step 4.
- 7 The recorder then finishes the calibration printout. Confirm that the line prints on the 20 unit gridline.

## Fetal Recorder Selftest Report

To verify your printer configuration, or if you doubt the performance of the recorder, you may want to print a test report.

To print a selftest report, in Service Mode, select **Main Setup -> Fetal Recorder-> Selftest**.

Here is an excerpt from a sample test report to give you an idea what it looks like (the exact appearance may vary slightly):



Same data from the trace header (see "Trace Header" on page 70)

Selftest status

Print quality data

Recorder electrical data

Paper-related data (Paper Count is the number of pages detected in this test)

SW Rev:	240	F.01.42
Serial Number:	09FB0542AF	
FPGA Rev:	B.01.05	
Protocol Rev:	A.01.05	
FPGA:	180	OK
EEPROM:	180	OK
Printhead/SCC:	OK	
Calibration:	50	OK
PH Offset:	2	
PS White:	120	
PS Black:	190	
PS Threshold:	70	
PS Intensity:	4	
PH Volt Min:	0	
PH Volt Max:	828	
PH Temperature:	396	
Motor Volt 1:	32	
Motor Volt 2:	84	
Speed Value:	7	
Paper Value:	2994	
Paper Count:	7	
Paper Type:	US (red)	

Example of selftest report

O <sub>2</sub>	40
PULSE	20
TEMP	0
B/P	0

This line should print exactly on the 20 unit gridline (Toco baseline) if the recorder is correctly calibrated.

Expected value for PH Volt Min is 0. If value is greater than 0, there may be a TLPH voltage problem.

Speed Value is the tolerance for paper detection. Value should be less than 20. If it is more, there could be a recorder assembly problem.

A Paper Value of 2994 is optimal: you may see small variations from this figure.

Check the test pattern to ensure all the heating elements on the printer head are operational. Ensure that:

- No more than 20 dots are missing over the entire printhead.
- No more than 2 adjacent dots are inoperative.

- No dots in the mode annotation (for example, FHR1) are inoperative.

If any of the above conditions are not met, replace the printhead (see “Removing the Thermal Line Printhead (TLPH)” on page 124).

Ensure that all printed lines are straight. If the lines are not straight, there may be a problem with the paper recorder speed.

## After Installation, Testing or Repair

Before handing the patient monitor over to the end-user, make sure it is configured appropriately and that it is in monitoring mode. Ensure that the user receives the current revision of the monitor documentation.



# Troubleshooting

A list of system error messages and troubleshooting information for common problems you may encounter while using the monitor and its accessories is given in the *Instructions for Use*. This chapter provides a guide for qualified service personnel for troubleshooting problems that cannot be resolved by the user.

**CAUTION** If the troubleshooting procedure requires you to disassemble the monitor or transducers, be certain to follow the disassembly and reassembly procedures given in Chapter 10, “Disassembly and Reassembly”.

## Who Should Perform Repairs

Only qualified service personnel should open the monitor housing, remove and replace components, or make adjustments. If your medical facility does not have qualified service personnel, contact Philips’ Response Center or your local Philips representative.

**WARNING** **High Voltage** - Voltages dangerous to life are present in the instrument when it is connected to the mains power supply. Do not perform any disassembly procedures with power applied to the instrument. Failure to adhere to this warning could cause serious injury or death.

## Replacement Level Supported

The replacement level supported for this product is to the printed circuit board (PCB) and major subassembly level. Once you isolate a suspected PCB, follow the procedures in Chapter 10, “Disassembly and Reassembly” to exchange the PCB with a known good replacement. Check to see if the symptom disappears and that the monitor passes all performance tests. If the symptom persists, swap back the replacement PCB with the suspected malfunctioning PCB (the original PCB that was installed when you started troubleshooting) and continue troubleshooting as directed in this chapter.

## Checking Revision Information

There are various ways to check revision information:

- Most of the revision information can be checked from reading the contents of the trace header (see “Trace Header” on page 70).
- You can also identify the hardware revision via the Main Setup menu (see “Hardware Revision Check” on page 70).

- You can also identify the software revision via the Main Setup menu (see “Software Revision Check” on page 71).

## Trace Header

The trace header printed when the recorder starts contains useful information about the monitor and its parameters.



- The first line (marked 1) contains:
  - the time.
  - date.
  - set paper speed.
- The second line (marked 2) contains:
  - result of the selftest.
  - firmware revision of the FPGA microcontroller, responsible for controlling the recorder, the display and the ultrasound tone. This revision is always fixed with a particular software revision.
  - revision of the internal recorder communication protocol. This revision is always fixed with a particular software revision.
  - paper type (US or INT).
  - information on whether the recording is a real-time trace (RT appears), or a stored data/trace recovery printout (RT is not printed).
  - information on the printhead intensity (I).
- The rest of the information (marked 3) contains:
  - product serial number and firmware revision.
  - Bus Master firmware revision (OB).
  - serial number and firmware revision of connected devices.

If a transducer is plugged into the monitor after the recorder started, the serial number and revision information is annotated along the bottom of the trace.

## Hardware Revision Check

Some troubleshooting tasks may require that you identify the hardware revision of your monitor’s main board. To check your hardware revision:

- 1 Enter the Main Setup menu and select **Revision**.
- 2 Select **Product**.

You see the hardware revision in the pop-up window, along with the serial number, part number, and the software revision.

## Software Revision Check

Some troubleshooting tasks may require that you identify the software revision of your monitor. You can find the software revision along with other information, such as the system serial number, in the monitor revision screen. To access the monitor revision screen:

- 1 Enter the Main Setup menu and select **Revision**.
- 2 Select **Product**.

You see the software revision in the pop-up window, along with the serial number, part number, and the hardware revision.

**NOTE** The part numbers listed in the monitor revision screen do not necessarily reflect the part numbers required for ordering parts. Please refer to Chapter 9, “Parts” for the ordering numbers. Photos of the parts are included for swift identification.

**NOTE** The system serial number can also be found on the rear of the monitor.

## Obtaining Replacement Parts

See Chapter 9, “Parts” section for details on replacement parts.

## Troubleshooting Guide

Problems with the monitor are separated into the categories indicated in the following sections and tables. Check for obvious problems first. If further troubleshooting instructions are required refer to the Troubleshooting Tables.

Taking the recommended actions discussed in this section will correct the majority of problems you may encounter. However, problems not covered here can be resolved by calling Philips Response Center or your local representative.

### Checks for Obvious Problems

When first troubleshooting the instrument, check for obvious problems by answering basic questions such as the following:

- 1 Is the power switch turned on?
- 2 Is the AC power cord connected to the instrument and plugged into an AC outlet?

### Checks Before Opening the Instrument

You can isolate many problems by observing indicators on the instrument before it is necessary to open the instrument.

### Checks with the Instrument Switched On, AC connected

The green power LED lights for about 1.5 seconds after switching on, and then goes out, and remains unlit during normal operation. The location of the green LED is shown in the following photograph:



## Individual Parameter INOPs

If you see any of the following parameter INOPs:

DECG EQUIP MALF	IUP EQUIP MALF
ECG EQUIP MALF	NBP EQUIP MALF
FetRec EQUIP MALF	OB EQUIP MALF
FHR1 EQUIP MALF	SpO <sub>2</sub> EQUIP MALF
FHR2 EQUIP MALF	SpO <sub>2</sub> SENSOR MALF
FHR3 EQUIP MALF	TOCO EQUIP MALF

try exchanging the relevant component (transducer, sensor, patient module or board) with a known good replacement, following the procedures in Chapter 10, “Disassembly and Reassembly”. Check to see if the INOP disappears, and that you can measure the parameter in question normally. If the INOP persists, swap back the original component and continue troubleshooting as directed in this chapter.

If you see the **OB EQUIP MALF** INOP following a monitor software upgrade, it is likely that the firmware in the bus master board is incompatible with the new software. Check the firmware revision, and upgrade this if necessary with the Support Tool. Contact Philips Support for more information regarding software and firmware revisions.

After checking/upgrading the bus master firmware, and you still suspect a defective bus master board, first try plugging the transducers into another monitor. If the transducers work properly with the other monitor, then exchange the bus master board.

In the case of the INOPs **FHR1 EQUIP MALF**, **FHR2 EQUIP MALF**, and **FHR3 EQUIP MALF**, when there are two or more ultrasound transducers attached to the monitor, identify the transducer for which the INOP was issued, using the blue transducer Finder LED. Touching a numeric on the screen makes the Finder LED light on the transducer providing the measurement. If you cannot identify the suspected transducer directly because the transducer Finder LED does not light due to the defect, identify the other, functioning transducers by activating their Finder LEDs, thus finding the defective one by a process of elimination.

## Initial Instrument Boot Phase

The following table describe the regular initial boot phase of the monitor. If the boot phase does not proceed as described below go to Boot Phase Failures for Troubleshooting information.

Time (sec.) after Power On	Boot Phase Event
0	Switch the monitor on using the On/Off switch.
2	The green AC Power LED lights for about 1.5 seconds.
3.5	Green AC Power LED is turned off, and remains off.
5	You hear a 'pop' from the loudspeaker
6-8	Boot Screen with the Philips Logo appears on the display. Test Sound is issued.
8-10	Boot Screen with the Philips Logo disappears Fixed screen elements (for example smart keys, alarm fields) appear on the screen.
10-15	First measurement information appears on the screen, touchscreen is functional.

## Troubleshooting Tables

The following tables list troubleshooting activities sorted according to symptoms.

### How to Use the Troubleshooting Tables

The possible causes of failure and the remedies listed in the troubleshooting tables should be checked and performed in the order they appear in the tables. Always move on to the next symptom until the problem is solved.

Boot Phase Failures

Screen is Blank

Touchscreen Not Functioning

General Monitor INOP Messages

Alarm Tones

Fetal Recorder

LAN / RS232

## Boot Phase Failures

Symptoms	Possible Causes of Failure	Failure Isolation and Remedy
Green LED does not light up, and no test tone is heard	No AC mains connection	Check that the power cord is not damaged and is properly connected to the monitor. Check that the power cord is correctly connected to a powered AC mains socket.
	Power supply defective	Remove power supply and check if output voltage is within the specifications (24V). Measure on multi-colored wired connection between red and black wires Exchange power supply if defective
	Power On/Off switch defective	Replace power supply
	Aborted/interrupted or inconsistent software configuration	Perform a software upgrade using the Support Tool.
	Main CPU Board defective	Replace Main CPU Board. Add boards in reverse order and try again with each board.
Green LED does not light up, but you hear a test tone	Display Assembly not connected to the Main CPU Board	Check if Display Assembly is connected correctly to the Recorder Adapter Board. Check that the multi-pin connector between the Recorder Adapter Board and the Main CPU Board
	Touch controller defective	Replace the Display Assembly.
	Display Adapter Board defective	Disconnect and reconnect the flat cable of the Display Adapter Board and check again
	LED defective	Try to switch on the monitor. If it operates normally, the LED is defective. Repair is effected by replacing the Display Assembly.
Green LED stays on continuously	Main CPU Board defective	Try loading new software. If this does not solve the problem, replace Main CPU Board.
Green LED blinks (indicating cyclic reboots)	Hardware failure	Connect Support Tool directly to monitor with crossover cable and start "search for defective devices" If no device is detected, proceed as described above in the section "Green LED stays on continuously"
	Software fault	If the Support Tool can detect the device and it indicates the Operating Mode is 'Boot', download and store the status log. Reload software and re-clone the monitor. If this fixes the problem e-mail the status log to your local response center
	Hardware failure	If this does not rectify the problem, follow the instructions under "Green LED stays on continuously".

Symptoms	Possible Causes of Failure	Failure Isolation and Remedy
No Test Sound issued or <b>INOP Speaker Malfunct.</b> issued	Speaker cable disconnected	Check speaker connections.
	Speaker defective	Check for INOPs and follow instructions Exchange speaker
	Main CPU Board defective	Exchange Main CPU Board

## Screen is Blank

The information listed in this table is only valid if the boot phase has completed without error. See Boot Phase Failures table for a description of the boot phase.

Symptoms	Possible Causes of Failure	Failure Isolation and Remedy
Display is blank or brightness is reduced	Display Adapter Board cable not connected	Check cable connection of Display Assembly to the Recorder Adapter Board.
	Backlight tubes defective	Replace Display Assembly.
	Backlight inverter defective	
	Display adapter board defective	
	LCD flat panel defective	
	Main CPU Board defective	Replace Main CPU Board.

## Touchscreen Not Functioning

Symptoms	Possible Causes of Failure	Failure Isolation and Remedy
Touchscreen not functioning	Touchscreen functionality has been temporarily disabled	Check if touchscreen functionality has been temporarily disabled (padlock symbol on Main Screen key). If yes, press and hold the Main Screen key to re-enable touchscreen operation.
	Touch screen cable not connected	Check connection from the Display Assembly to the Recorder Adapter Board. If the problem is not resolved, check that the multi-pin connector between the Recorder Adapter Board and the main CPU Board.
	Touch controller defective	Replace Display Assembly
	Touch Sensor defective	
	Main CPU Board defective	Replace Main CPU Board
Touch Position invalid	Touch not calibrated	Perform touch calibration: 1 Enter Service Mode 2 Enter the Main Setup Menu 3 Select <b>Hardware</b> 4 Select <b>Calibrate Touch</b> See “Touchscreen Calibration” on page 64

## General Monitor INOP Messages

INOP Message	Possible Causes of Failure	Failure Isolation and Remedy
<b>CheckInternVoltage</b>	Problem with the voltages (5V) in the monitor	Remove all I/O boards and put them back in one at a time to isolate any defective board. If this does not resolve the problem, replace the main board.
<b>Check Monitor Func</b>	The temperature inside the monitor is too high	Check the environment for possible causes.
	Main Board defective	Replace Main Board.

INOP Message	Possible Causes of Failure	Failure Isolation and Remedy
<b>Check Settings</b>	INOP occurs during normal operation, indicating a possible monitor software problem	<p>Check the monitor and patient settings before you resume monitoring. If the settings are unexpected, there may be a problem with the monitor software.</p> <ol style="list-style-type: none"> <li>1 Silence the INOP.</li> <li>2 Load the User Defaults (see “Loading the User Defaults” on page 164).</li> <li>3 If this is unsuccessful, try loading the Factory Default (see “Loading the Factory Default” on page 164), and reconfigure the monitor in Configuration Mode, and save the new settings in the User Defaults.</li> </ol> <p>If the INOP persists, there is an unresolved software problem. Report the problem to factory support.</p>
	INOP occurs after a software upgrade, indicating a possible incomplete or unsuccessful upgrade	Clone the correct settings via the Support Tool.
<b>Internal.Comm.Malf.</b>	Main CPU Board defective	Replace Main CPU Board.
<b>Settings Malfunc.</b>	Problem during cloning process.	Reclone configuration file.
	Memory space in which the settings are stored has been corrupted	Reclone configuration file. This will reload the memory space.
	Main CPU Board defective	Replace Main CPU Board.

## Keyboard/Mouse Not Functioning

Symptoms	Possible Causes of Failure	Failure Isolation and Remedy
Keyboard/Mouse attached directly to the monitor not functioning	Keyboard/Mouse not connected properly	Check cabling
	Keyboard/Mouse defective	Replace Keyboard/Mouse
	PS/2 I/O board is not properly plugged in	Ensure the PS/2 I/O board is properly plugged in. If necessary, remove the board and plug it in again.
	PS/2 I/O board defective	Replace I/O board

## Network Status Icons

Icon	Explanation
No Icon	LAN cable not connected (monitor does not have a LAN connection).
	LAN cable connected, no connection to OB TraceVue. To check whether an IP Address has been assigned, enter <b>Main Setup --&gt; Bed Information</b> and scroll to <b>IP Address</b> . (0.0.0.0 means no IP address has been assigned.) OB TraceVue may send a prompt message, giving the reason why connection to OB TraceVue cannot take place.
	LAN cable connected, IP address assigned, monitor connected to OB TraceVue. OB TraceVue may send a prompt message, indicating a possible problem.

## Alarm Tones

Symptoms	Possible Causes of Failure	Failure Isolation and Remedy
INOP Message <b>Speaker Malfunc.</b> is displayed	Speaker cable disconnected	Reconnect speaker cable
	Speaker defective	Replace speaker
	Sound amplifier on Main CPU Board defective	Main CPU Board
Alarm occurs but no alarm sound is issued	Volume set to 0	Increase volume
	Speaker defective	Replace speaker
	Sound amplifier on Main CPU Board defective	Main CPU Board

## Alarm Behavior

If your monitor did not alarm in the way in which the end user expected, please consult the *Instructions for Use* for possible setup issues or configuration settings which could affect alarm behavior.

## Fetal Recorder

Symptom	Possible Cause	Corrective Action
Paper empty warning is issued in the status line at the bottom of the screen, but paper is not out.	Drawer is open.	Close the drawer.
	Paper jam.	Open the drawer, remove paper, tear off scrumpled paper and re-load, or load a new pack of paper. Close the drawer.
	Paper sensor dirty.	Clean paper sensor (see Chapter 7, “Testing and Maintenance”).
	Paper sensor defective.	Exchange paper sensor (see Chapter 10, “Disassembly and Reassembly” and Chapter 9, “Parts”).

Symptom	Possible Cause	Corrective Action
No paper transport.	Poor connection.	Check all internal connectors.
	Paper jam.	Open the drawer, remove paper, tear off scrumpled section of paper and re-load, or load a new pack of paper. Close the drawer.
	Motor cable is disconnected.	Check that the motor cable is properly connected to the Recorder Adapter Board.
	Motor is defective.	To test the functioning of the motor, open the drawer and press the recorder <b>Start/Stop</b> key to start the recorder. A good motor should rotate for between one and three minutes (depending on the paper speed).  If the motor does not rotate, replace the motor (see “Replacing the Stepper Motor” on page 131).
	Drawer is open.	Close the drawer.
The recorder appears to be running normally, but the paper remains blank	Thermal Printhead is disconnected.	Check the connection. Then run the recorder Selftest to verify correct printing (see “Fetal Recorder Selftest Report” on page 66).
	Thermal Printhead is defective.	Replace the Thermal Printhead. Then calibrate the recorder (see “Setting the Fetal Recorder Offset” on page 65).
	The wrong side of the paper is facing up.	Load the paper correctly, the right way up.
No recorder key is available on the screen, and the <b>INOP FetRec EQUIP MALF</b> is issued.	The recorder has not been calibrated.	Calibrate the recorder (see “Setting the Fetal Recorder Offset” on page 65).
	EEPROM on the Recorder Adapter Board is defective	Exchange the Recorder Adapter Board and calibrate the recorder (see “Setting the Fetal Recorder Offset” on page 65).
	Recorder Controller on the Main CPU Board is defective.	Exchange the Main CPU Board and calibrate the recorder (see “Setting the Fetal Recorder Offset” on page 65).
	Recorder cable is disconnected.	Ensure the recorder cable is connected firmly at both ends.

Symptom	Possible Cause	Corrective Action
The INOP <b>CHECK PAPER</b> is issued.	The drawer is open and there is paper on the paper sensor.	Ensure the paper is loaded correctly, and close the drawer.
	Paper jam.	Open the drawer, remove paper, tear off scrumpled section of paper and re-load, or load a new pack of paper. Close the drawer.
	Paper sensor dirty.	Clean paper sensor (see Chapter 7, "Testing and Maintenance").
	Paper sensor defective.	Exchange paper sensor (see Chapter 10, "Disassembly and Reassembly" and Chapter 9, "Parts").
	The rubber roller is dirty.	Clean the rubber roller (see Chapter 7, "Testing and Maintenance").
	Paper is not approved by Philips.	Use only paper approved by Philips.
	Inadequate contrast of paper marks.	Use only Philips approved paper. Calibrate the recorder.
The INOP <b>WRONG PAPER SCALE</b> is issued.	Paper with the wrong scale has been loaded (for example, International paper has been loaded instead of US paper).	Check, and if necessary, replace the paper pack with one with the correct scale. Check, and if necessary, change the paper scale setting to the correct setting for the paper used.
The INOP <b>PRINthead OVERHEAT</b> is issued.	The printhead is too hot.	Wait for the printhead to cool down, then press the recorder <b>Start/Stop</b> key or the <b>Silence</b> key to clear the INOP.
Bad or distorted printout within the first 1 cm of the trace.	Paper drawer was not fully closed.	Always ensure the paper drawer is fully closed before starting recording.
Poor print quality.	Heat setting needs adjusting.	Adjust the Thermal Printhead heat setting. Then run the recorder Selftest to verify correct printing (see "Fetal Recorder Selftest Report" on page 66).
	Thermal Printhead dirty.	Clean the Thermal Printhead (see Chapter 7, "Testing and Maintenance"). Then run the recorder Selftest to verify correct printing (see "Fetal Recorder Selftest Report" on page 66).
	Thermal Printhead failure.	Exchange the Thermal Printhead (see "Removing the Thermal Line Printhead (TLPH)" on page 124 and "Replacing the TLPH" on page 125). Then run the recorder Selftest to verify correct printing (see "Fetal Recorder Selftest Report" on page 66).
Paper not feeding properly.	Paper incorrectly loaded.	Load paper correctly.
	The rubber roller is dirty.	Clean the rubber roller (see Chapter 7, "Testing and Maintenance").

Symptom	Possible Cause	Corrective Action
Trace is not printed correctly with reference to the paper gridlines.	Offset needs adjusting.	Calibrate the recorder and change the offset (see “Setting the Fetal Recorder Offset” on page 65).
	The wrong paper scale is being used.	Ensure the paper you are using matches the paper scale setting.

## LAN / RS232

Symptoms	Cause of Failure	Failure Isolation and Remedy
External device (such as a surveillance system like OB TraceVue) not receiving data	The LAN/RS232 port is not configured for data export	Check configuration of the LAN/RS232 ports in configuration mode
	The cable between the external device and the monitor is not connected correctly or defective	Check cable and replace if necessary
	The external device does not support the version of the data export protocol used in the monitor	Check if the device supports the version of the data export protocol. Upgrade device or monitor if necessary (if matching versions exist).
	A terminal concentrator is used in between the device and the monitor and a protocol with dynamic speed negotiation is used	Some terminal concentrators do not support changing the transmission speed (baud rate) dynamically. Check if the connection works without the concentrator
	The LAN/RS232 board is in a wrong slot (slot has been changed after software configuration or an additional board has been plugged in)	Verify correct placement of the I/O boards
	The LAN/RS232 board is defective	Check board and replace if necessary

## Transducers

Note that immediately after plugging in a normally functioning transducer, the Finder LED briefly lights twice.

Symptoms	Possible Cause	Failure Isolation and Remedy
<p>The transducer Finder LED does not light up after plugging in the transducer, and the transducer appears not to work.</p> <p><b>INOP OB EQUIP MALF</b> is displayed.</p>	Defective transducer cable.	Visually inspect the transducer cable and the cable connector for damage. If there are obvious signs of damage, replace the cable.
	Defective connector block.	Visually inspect the connector block and the sensor sockets for damage. If there are obvious signs of damage, replace the connector block.
	Transducer or connector block is defective.	<p>Try plugging the transducer into a different sensor socket.</p> <ul style="list-style-type: none"> <li>If the Finder LED works, then the original socket is defective. Replace the connector block.</li> <li>If the Finder LED still does not light in any of the other sockets, try using a known good transducer. If the Finder LED lights, the original transducer is defective: replace it.</li> </ul>
	Bus Master Board is defective.	Try using a known good transducer. If the Finder LED does not light in any of the sockets using a known good transducer, then the Bus Master Board is defective. Replace the Bus Master Board.
	No power to Bus Master Board.	If both the SpO <sub>2</sub> board and the Bus Master Board are not working, exchange the power supply.
<p>Transducer appears not to work, and:</p> <p><b>EITHER</b> the transducer Finder LED lights briefly after plugging in the transducer and INOP OB EQUIP MALF is displayed.</p> <p><b>OR</b> the transducer Finder LED lights briefly after plugging in the transducer, and there are no parameters on the screen (transducer is not recognized).</p> <p><b>INOP OB EQUIP MALF</b> is displayed.</p>	Transducer is defective.	Replace transducer.
All transducers (US, Toco, IUP and ECG) do not work.  <b>INOP OB EQUIP MALF</b> is displayed.	Bus Master Board is defective.	Replace Bus Master Board.

Symptoms	Possible Cause	Failure Isolation and Remedy
Transducer is connected, <b>INOP OB EQUIP MALF</b> is displayed.	Main CPU board is defective.	Replace Main CPU Board.
	Transducer defective.	Replace transducer.
	Interrupted transducer upgrade, or software based malfunction of the transducer, including communication problems between the Bus Master Board and the transducer.	Perform software upgrade of the transducer with the Support Tool.
Transducer belt button is broken or damaged.	Mechanical damage.	Replace the belt button. Handle transducers with care. <b>Never use a transducer with a broken or damaged knob.</b>

## Status Log

Many events that occur during start-up or regular monitoring are logged in the Status Log. The Status Log can be cleared. Not all entries in the Status Log are errors. You can print the Status Log via the Support Tool.

Monitor Id.	Code	No.	Date	
			Time	
H	18202	20100	1	4 Apr 05 16:37
C	1721	21050	1	4 Apr 05 15:37

The Status Log window shows logged events which caused a reboot of the monitor.

To enter the Status Log Window, select Main Setup -> Revision. The following list opens up:

- **Status Log**
- **Product**
- **Appl. SW**
- **Config**
- **Boot**
- **Language**
- **Settings**
- **OB**
- **FetRec**

- **NBP** (optional)
- **SpO<sub>2</sub>** (optional)
- **List of plugged parameters**

Select **Status Log**.

The first column in the log identifies the event class (“C”: caused a cold start, “H”: caused a hot start, “N”: no restart, for information only). Column 3 and 4 identify the event source and event code. Column 4 counts the number of occurrences of the event. The last column shows the time and date of the last occurrence of the event.

The following pop-up keys overlay the SmartKeys:

<b>Clear StatLog</b>		<b>M2703A</b>	
----------------------	--	---------------	--

#### **Clear StatLog**

This key clears the currently displayed Status Log

#### **M2703A**

This key switches to the Monitor Revision Window

If an event occurs repeatedly, contact your Philips Service Representative.

**NOTE** It is possible, using the support tool, to download the status log and send it to your Philips Service Representative as a file (for example via e-mail).

## **Troubleshooting with the Support Tool**

Using the support tool you can:

- access the full status log which can be saved as a file
- reload software
- identify defective devices
- start touch screen calibration

For details on how to perform these tasks see the Support Tool User Manual.

## **Troubleshooting the Individual Measurements or Applications**

For problems isolated to an individual parameter or application, please consult the *Instructions for Use* and configuration information.

If the *Instructions for Use* did not resolve an individual parameter problem, then another transducer or patient module should be tried.

If you are getting questionable readings for individual measurements you may want to do the performance assurance tests in Chapter 7, “Testing and Maintenance”.

The performance of the individual applications are affected by the configuration of the monitor. When contacting Philips support you may be asked about the configuration of the monitor to aid in troubleshooting.



# Parts

Spare parts, along with part numbers, are listed in the tables that follow.

## Monitor

Description	Ordering Number				Qty	
	Number for New Parts		Number for Exchange Parts			
	Part No.	Alternative Identifier	Part No.	Alternative Identifier		
Cover, Connector Symbol without SpO <sub>2</sub> and Noninvasive Blood Pressure	M2703-44103	4512 610 10311	-	-	1	
Cover, Noninvasive Blood Pressure Symbol	M2703-44105	4512 610 10321	-	-	1	
Cover, all inc. SpO <sub>2</sub> and Noninvasive Blood Pressure Symbol	M2703-44106	4512 610 10331	-	-	1	
Cover, Connector Text without SpO <sub>2</sub> and Noninvasive Blood Pressure	M2703-44113	4512 610 10341	-	-	1	
Cover, Connector Noninvasive Blood Pressure Text	M2703-44115	4512 610 10351	-	-	1	
Cover, Connector inc. SpO <sub>2</sub> and Noninvasive Blood Pressure Text	M2703-44116	4512 610 10361	-	-	1	
Power Supply Assembly	M2703-68001	4512 610 07271	-	-	1	
Loudspeaker Assembly	M2703-60002	4512 610 10231	-	-	1	
Paper Sensor Assembly	M2703-60003	4512 610 10411	-	-	1	
Stepper Motor Assembly	M2703-60004	4512 610 10401	-	-	1	
Bottom Housing Assembly	M2703-64101	4512 610 10221	-	-	1	
Top Cover Housing	M2703-64102	4512 610 10391	-	-	1	
Top Cover Assembly	M2703-60502	4512 610 11201	-	-	1	
Display Assembly (see page 97 for assembly contents)	M2703-64503	4512 610 10441	M2703-68503	4512 610 11221	1	

<b>Description</b>	<b>Ordering Number</b>				<b>Qty</b>	
	<b>Number for New Parts</b>		<b>Number for Exchange Parts</b>			
	<b>Part No.</b>	<b>Alternative Identifier</b>	<b>Part No.</b>	<b>Alternative Identifier</b>		
Noninvasive Blood Pressure Assembly (picture on page 94)	M2703-64502	4512 610 10271	M2703-68502	4512 610 10551	1	
Housing Connector Noninvasive Blood Pressure	1253-8416	4512 610 10281	-	-	1	
SpO <sub>2</sub> Board	M1020-66513	4512 610 16351	-	-	1	
Housing Connector SpO <sub>2</sub>	1253-8422	4512 610 10301	-	-	1	
Paper Drawer Assembly	M2703-64651	4512 610 10431	-	-	1	
Main CPU Board	M2705-66510	451261024951	M2705-68510	451261024961	1	
Bus Master Board	M2703-66520	4512 610 11191	-	-	1	
Socket Connector Block	1253-8415	4512 610 10261	-	-	1	
Recorder Adapter Board	M2703-66530	4512 610 11211	-	-	1	
Thermal Line Printhead	1810-2440	4512 610 10381	-	-	1	
LAN / RS232 Interface Assembly	M2703-67501	4512 610 10531	-	-	1	
Input Device Interface (2x PS/2)	M8086-67501	4535 634 69651	-	-	1	
Lever Stop (pack of 5)	M2703-64205	4512 610 10521	-	-	1	
FM Small Parts Kit - Plastic Parts and Labels (see page 99 for kit contents)	M2705-64203	4512 610 25031	-	-	1	
FM Small Parts Kit - Screws and Cables (see page 100 for kit contents)	M2703-64202	4512 610 25041	-	-	1	

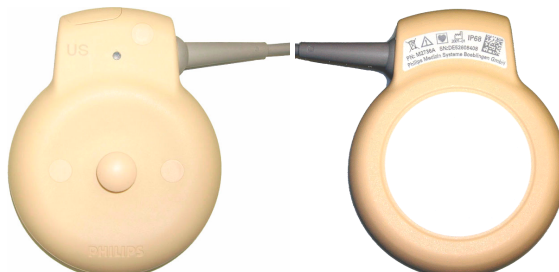
# Transducers



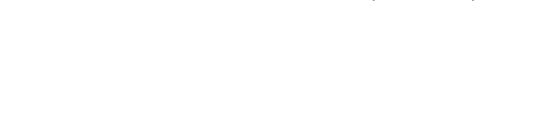
Toco Transducer (M2734A)



Toco<sup>+</sup> Transducer with ECG/IUP capability  
(M2735A)



Ultrasound Transducer (M2736A)



Description	Ordering Number				Qty	
	Number for New Parts		Number for Exchange Parts			
	Part No.	Alternative Identifier	Part No.	Alternative Identifier		
Toco Transducer	M2734-60501	451261010451	M2734-68501	4512 610 11231	1	
Toco+ Transducer	M2735-60501	451261010461	M2735-68501	4512 610 11241	1	
US Transducer	M2736-60501	451261010471	M2736-68501	4512 610 11251	1	
Cable Assembly (for all transducers; see page 98)	M2735-64201	4512610 0481	-	-	1	
Belt Button Kit, with tool, pack of 5 (see page 98 for kit contents)	M2703-64204	451261010511	-	-	1	

# Patient Modules



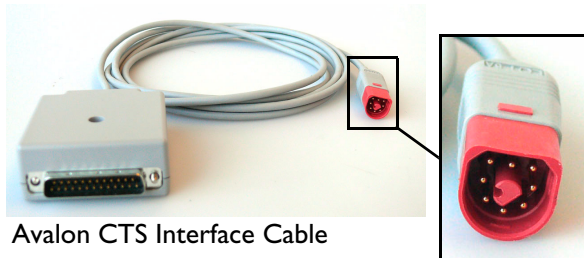
Patient module for ECG/IUP  
(M2738A)



Remote Event Marker  
(989803143411)

Description	Ordering Number				Qty	
	Number for New Parts		Number for Exchange Parts			
	Part No.	Alternative Identifier	Part No.	Alternative Identifier		
ECG/IUP Patient Module	M2738-60501	451261011261	-	-	1	
Remote Event Marker	-	989803143411	-	-	1	

# Interface Cables



Avalon CTS Interface Cable  
(M2731-60001) for front  
connection

Description	Ordering Number				Qty	
	Number for New Parts		Number for Exchange Parts			
	Part No.	Alternative Identifier	Part No.	Alternative Identifier		
Avalon CTS Interface Cable (M2731-60001), red connector, for connection to the red fetal sensor sockets.	M2731-60501	451261016251	-	-	1	
Serial (RS232) Interface Cable to OB Tracevue.	M1380-61612	453563278111	-	-	1	

# Mounting Hardware

Description	Product Option Number	Ordering Number		Qty
		Part No.	Alternative Identifier	
Flush Wall Mount for flat wall mounting)	M2740A #A01	M2740-64001	4512 610 09061	1
Wall Mounting Arm with tray	M2740A #A05	M2740-64002	4512 610 09071	1
Cart with fixed angle mount and two drawers	M2740A #C01	M2740-64003	4512 610 09081	1
Roll Stand with tray	M2740A #R01	M2740-64005	4512 610 09101	1
Mounting Kit for Avalon CTS for use with M2740A #C01 cart, with complete system power from one single power cord, and integrated interface cable	M2740A #U01	M2740-64006	4512 610 16261	1
Wall Channel required for mounting of wall mounts (options A01 and A05)	M2740A #W01	5061-8324	4512 610 09111	1

M2740A #A01	M2740A #A05	M2740A #C01	M2740A #R01	M2740A #U01	M2740A #W01
					

**Flush Wall Mount** for flat wall mounting  
**Wall Mount (Arm)**  
**Cart** with fixed angle mount and two drawers  
**Rollstand** with tray  
**Mounting Kit** for Avalon CTS for use with M2740A #C01  
**Wall Channel** required for wall mounts A01 and A05

# Assembly and Kit Contents

The tables in this section provide additional information by listing the contents of assemblies and kits. Assemblies come fully assembled: the contents list details what is contained in the assembly.

## Bottom Housing Assembly



Bottom Housing Assembly Contents		Qty
Sub-Assembly	Contents	
Bottom Housing Assembly	Housing, Bottom	1
	Feet	1
Model/Serial Number Plate		1
Support Sub-Assembly	Support	1
	Pin, DIN 6325, 2.5 x 8	2
	Hinge Support	2
	Bracket	2
	Screw M3 x 6	4
	Catch, I/O Board	2
	Cable Holder	2
	Pin for Cable Holder	2

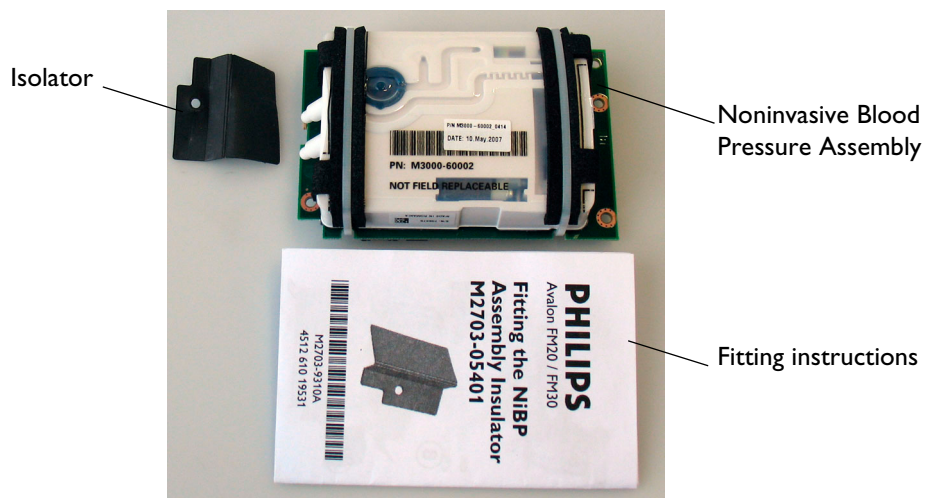
<b>Bottom Housing Assembly Contents</b>		<b>Qty</b>
<b>Sub-Assembly</b>	<b>Contents</b>	
Main Chassis Sub-Assembly	Chassis, sheet metal	1
	Standoff, M3 x 18	2
	Standoff, M3 x 10	2
	Standoff, M3 x 6	2
	Press Nut, M3	2
	Clip, RFI	4
	Cover, Board Guide	1
	Guide, I/O mid upper	3
	Guide, I/O mid lower	3
	RFI-Clip	2
	Screw, Torx, with washer, M3 x 6	5
	Holder, Loudspeaker	1
	Screw, Loudspeaker Holder	2

## Power Supply Assembly



<b>Power Supply Assembly Contents</b>	<b>Qty</b>
Power Supply Angle	1
Power Supply Frame	1
Nut, press in M3	3
Insulation	1
Screw with washer, M3 x 6	3

## Noninvasive Blood Pressure Assembly

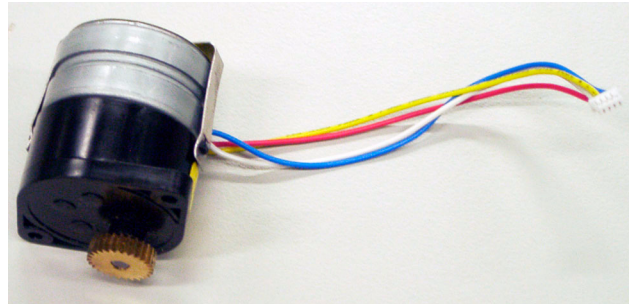


## Top Cover Housing



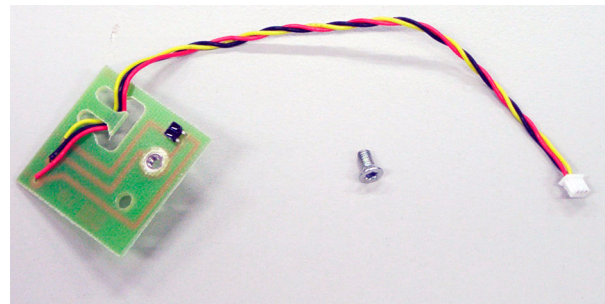
Top Cover Housing Contents	Qty
Top Cover	1
Handle	1
Paper Drawer Runner	2
Runner End-stop (left)	1
Runner End-stop (right)	1
Screw, Torx M3 x 8	4
Leaf Spring	1
Label, Avalon FM20	1
Label, Avalon FM30	1

## Stepper Motor Assembly



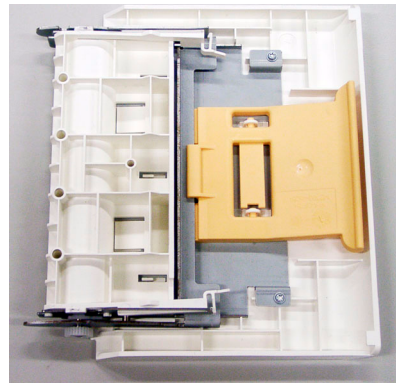
Stepper Motor Assembly Contents	Qty
Stepper Motor	1
Connector Housing	1
Connector Contact	1
Gearbox	1
Pinion	1

## Paper Sensor Assembly



Paper Sensor Assembly Contents	Qty
Paper Sensor, including cable and connectors	1
Nut, press-in M3	1
Screw, Torx M3 x 6	1

## Drawer Assembly



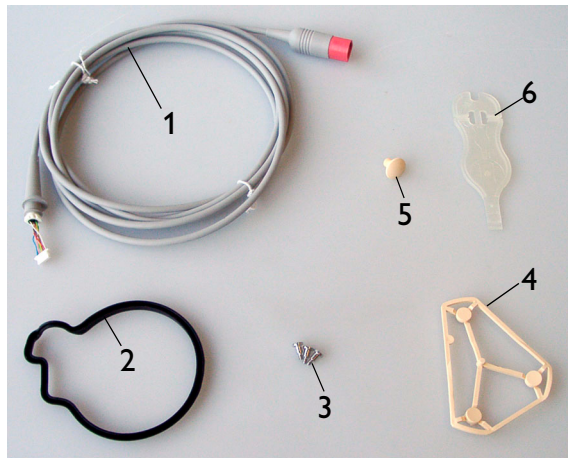
Drawer Assembly Contents	Qty
Paper Drawer Cover	1
Rubber Roller (including bearing, rod, pinion)	1
Platen Holder	1
Chassis Guide	1
Lever Stop	1
Latch	1

## Display Assembly



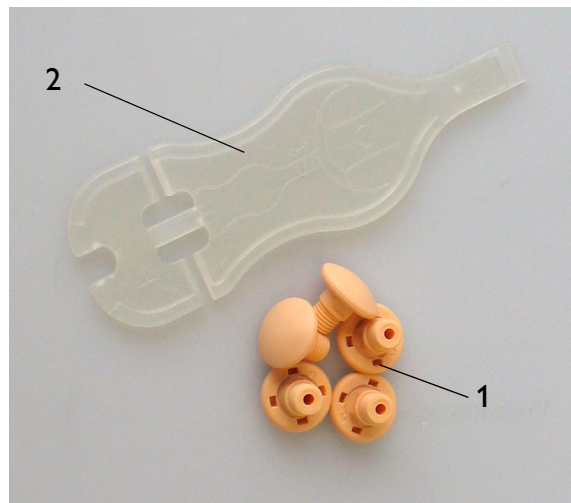
Display Assembly Contents	Qty
Housing, Bottom	1
Housing, Top	1
Clamp	1
Pin	2
Display Holder, lefthand	1
Display Holder, righthand	1
Backlight Tube	2
Hinge	2
Chassis Guide	1
Cable Guide, rear	1
Board Holder	5
Stop Lever	1
Ribbon Cable	1
PCA Touch Control	1
Inverter Board	1
TFT Display Unit	1
Touchscreen	1
Gasket (1050 mm)	1

## Transducer Cable Assembly



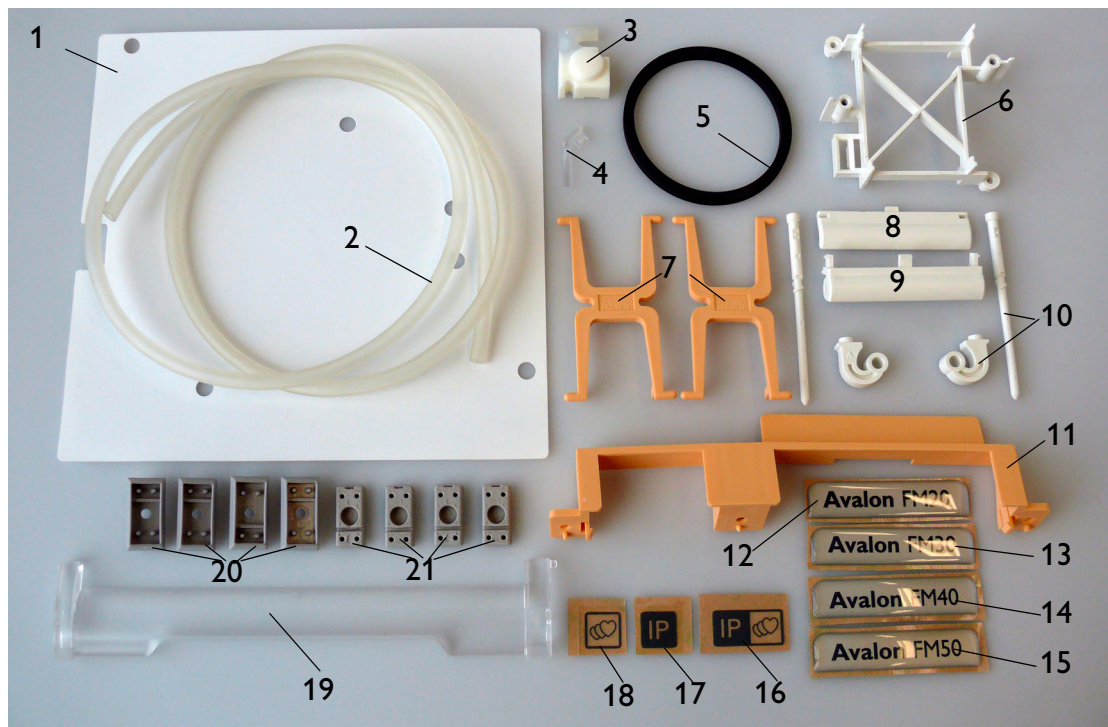
Item	Cable Assembly Contents	Qty
1	Transducer Cable (for all fetal transducers)	1
2	Sealing Gasket	1
3	Screw M2.5	3
4	Screw Cover (set of 3)	1
5	Transducer Belt Button	1
6	Avalon Tool	1

## Belt Button Kit



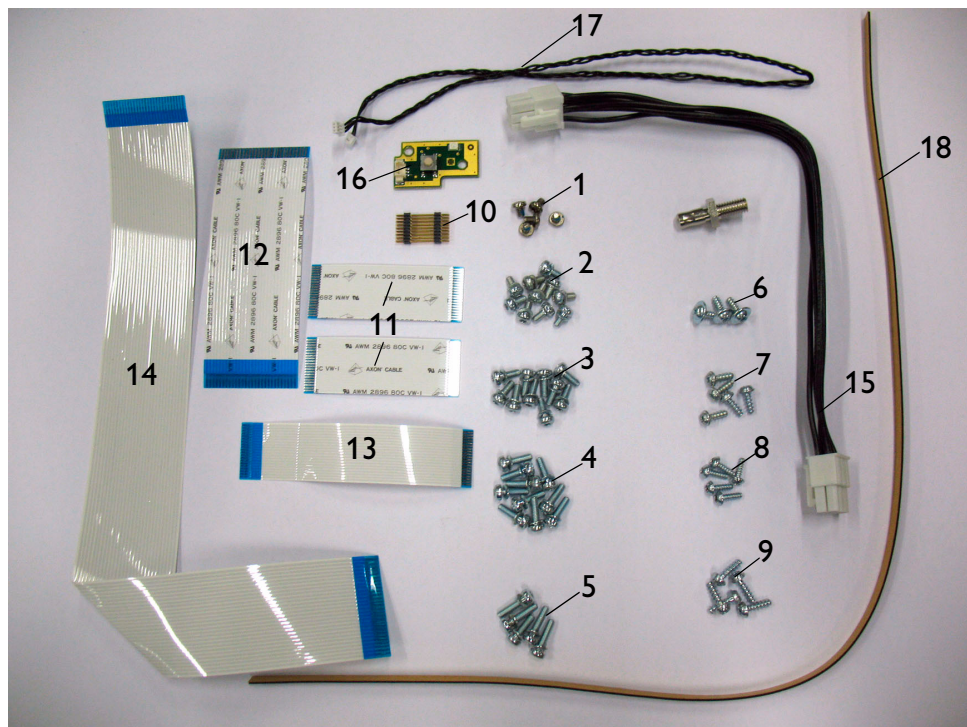
Item	Belt Button Kit Contents	Qty
1	Belt Buttons	5
2	Avalon Tool (for removing/replacing transducer belt buttons)	1

## FM Small Parts Kit - Plastic Parts and Labels



Item	FM Small Parts Kit Contents - Plastic Parts and Labels	Qty
1	FM40/50 Insulator, API Board	1
2	NBP Tubing (needs to be cut to size)	1
3	FM40/50 Silicone ON/OFF Key	1
4	FM40/50 Lightpipe	1
5	FM20/30 O-Ring, loudspeaker	1
6	FM40/50 Mounting Frame for SpO <sub>2</sub> Board	1
7	FM20/30 Ratchet Lever/Clip	2
8	FM20/30 Cable Guide, Rear	1
9	FM20/30 Cable Guide, Front	1
10	FM20/30 Hinge	2
11	FM20/30 Connector Frame	1
12	Label, FM20	1
13	Label, FM30	1
14	Label, FM40	1
15	Label, FM50	1
16	Intrapartum/Triplets Label	1
17	Intrapartum Label	1
18	Triplets Label	1
19	FM40/50 Paper Guide (incorporating tear-off edge)	1
20	FM40/50 Holder, Bumper Foot	4
21	FM40/50 Bumper Foot	4

## FM Small Parts Kit - Screws and Cables



Item	FM Small Parts Kit Contents - Screws and Cables	Qty
1	Torx M3 x 4	5
2	Torx M3 x 6 with washer	10
3	Torx M3 x 8 with washer	10
4	Torx M3 x 10 with washer	10
5	Torx M3 x 12 with washer	5
6	Screw Ejot K30 x 8	3
7	Screw Ejot 3 x 8	8
8	Screw Ejot 2.5 x 8	5
9	Screw Ejot 3 x 10	5
10	Connector for SpO <sub>2</sub> Board (all monitors)	1
11	Ribbon cable, Bus Master (all monitors)	2
12	FM20/30 Ribbon cable, Recorder	1
13	Ribbon cable, NBP to Main CPU Board (all monitors)	1
14	FM40/50 Ribbon cable, API Board to Recorder	1
15	FM40/50 Cable assembly, Power Supply	1
16	FM40/50 PCA, Switch Board	1
17	FM40/50 Switch Board cable to API Board	1
18	FM40/50 Sealing Gasket, Top Cover	1
19	FM40/50 Equipotential grounding bolt	1





# Disassembly and Reassembly

- 
- WARNING**
- Before attempting to open or disassemble the monitor, disconnect it from the AC mains supply.
  - Energized circuits are accessible with the covers open. Do not work on the monitor with the covers open and AC power connected. Only qualified service personnel should open or disassemble the monitor.
  - Performance verification: do not place the system into operation after repair or maintenance has been performed, until all performance tests and safety tests listed in Chapter 7, “Testing and Maintenance” have been performed. Failure to perform all tests could result in erroneous parameter readings, or patient/operator injury.
- 

**CAUTION** Observe ESD (electrostatic discharge) precautions when working within the unit.

---

## Introduction

Remember to store all screws and parts in a safe place for later refitting.

## How to Use this Chapter

The disassembly sections detail the step-by-step procedures you use to access replaceable parts of the monitor and the transducers.

The monitor consists of two major assemblies:

- The top cover assembly
- The bottom housing assembly

The top cover assembly consists of the top cover housing, the display assembly, the recorder assembly, and the recorder adapter board.

The bottom housing assembly consists of the bottom housing, the power supply assembly, the main CPU board, the bus master board, and depending on the options ordered, the noninvasive blood pressure assembly, the SpO<sub>2</sub> assembly, the input device interfaces, and the RS232/LAN interface.

All part numbers of spare parts are listed in Chapter 9.

## Tools Required

**CAUTION** When replacing the front cover, do not over-torque the screws. Excessive torque may damage the plastic screw mountings.

You need the following tools:



- Flat-head screwdriver, head thickness 0.5 mm to fit transducer screw
- Torx-head screwdriver, size T-10, minimum shaft length 80mm
- Small flat-head screwdriver, 2.0-3.0 mm
- Long-nosed pliers

## Screws Used

The following picture shows the range of screws used in the Avalon FM20/30/40/50 fetal monitors:



- Screw Ejot K30x8, T-10
- Screw Ejot 3x10, T-10
- Screw Ejot 3x8, T-10
- Screw Ejot 2.5x8, T-8
- Screw M3x12 with washer, T-10
- Screw M3x10 with washer, T-10
- Screw M3x8 with washer, T-10
- Screw M3x6 with washer, T-10
- Screw M3x4, T-10

# Serial Numbers

The serial number of the monitor appears on the device nameplate at the rear of the bottom housing. It is also stored electronically in the power supply.

- If you change the bottom housing, remove the nameplate from the old housing and fit it to the new housing.
- If you exchange the power supply of the monitor, you may have to re-enter the monitor serial number afterwards. Check the serial number of the monitor in the Support Tool device view to see whether this is necessary: if the sixth digit of a monitor serial number is an "X", you must re-enter the serial number, which you will find on the nameplate. Refer to the *Support Tool Instructions for Use* for details of how to change or re-enter a serial number.

# Top Cover Disassembly/Reassembly

This section describes disassembly and reassembly operations for the top cover and its assemblies.

## Removing the Top Cover Assembly

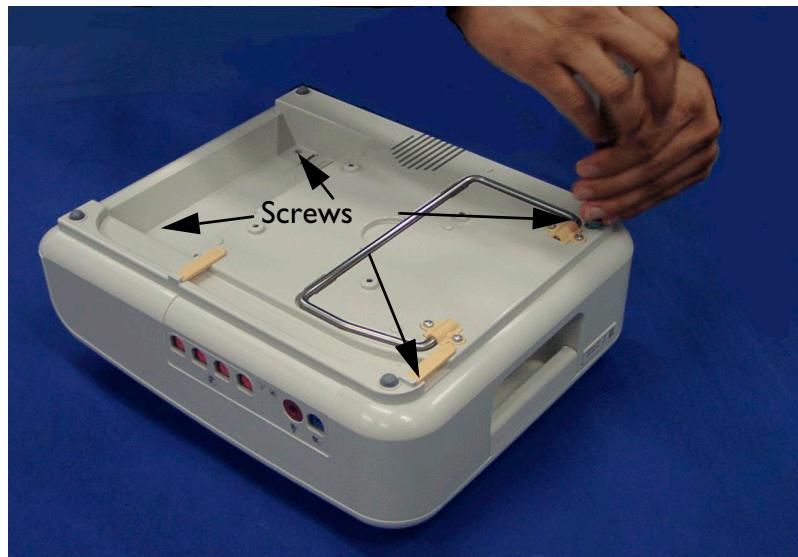
- 1 First fold the display completely flat.



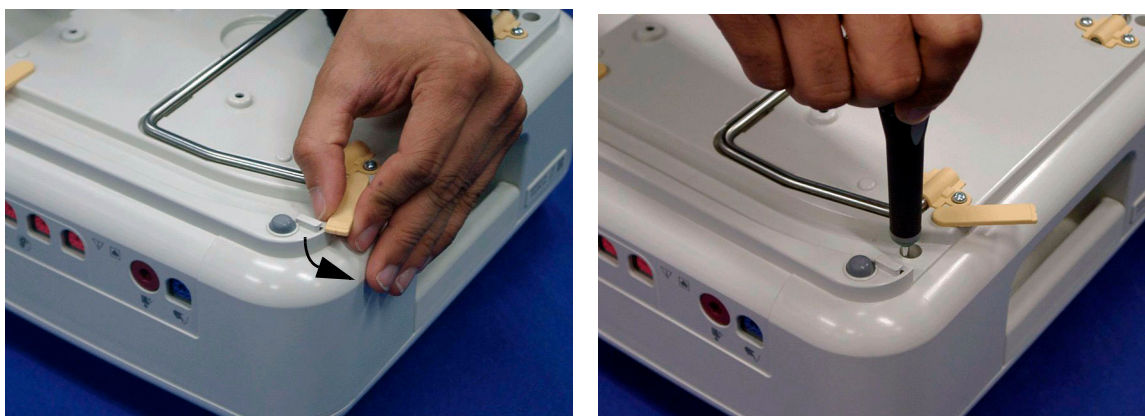
- 1 Carefully place the monitor upside down. To avoid scratches, place the unit on some cloth or other soft surface.



- 2 Remove the four screws securing the top cover assembly to the bottom housing, using a T-10 Torx driver.



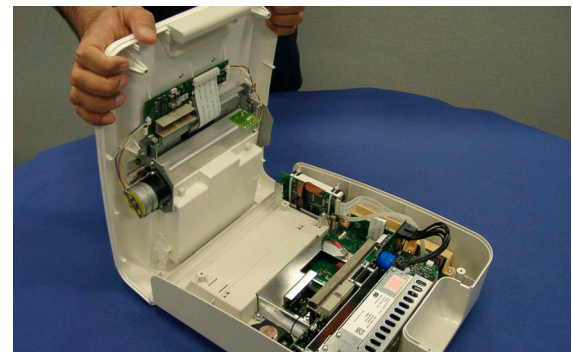
- 3 To gain access to the screw in the rear right hand corner, release the cable guide aside as illustrated.



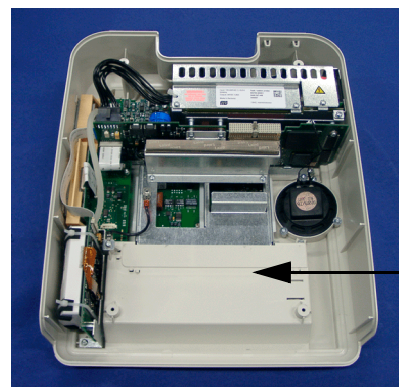
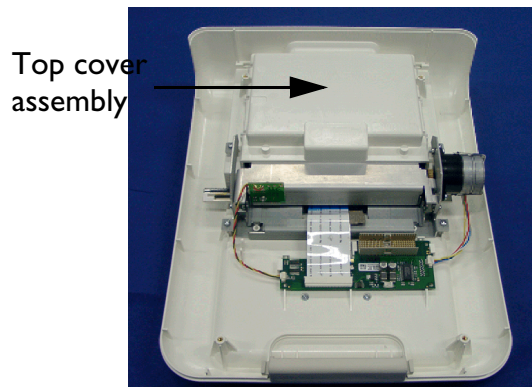
- 4 Holding both top cover and bottom housing assemblies together, place the monitor upright again.



- 5 Separate the top cover from the bottom housing from the front of the monitor as illustrated.



- 6 The top cover assembly is now separated from the bottom housing assembly.



## Removing the Display Assembly

The display assembly can be exchanged without removing the top cover of the monitor. Therefore, no additional safety tests are required after exchanging the display assembly.

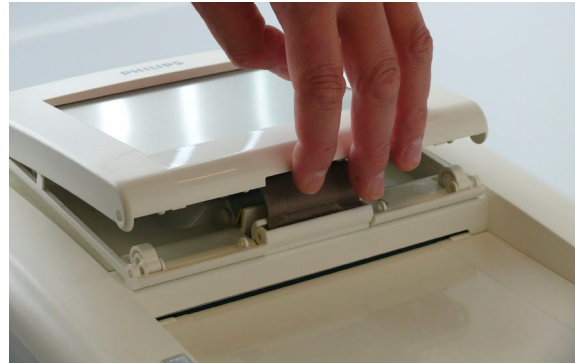
- 1 Squeeze the arms of the ratchet clip to remove it from the slots on the rear of the display housing.



- 2 Remove the two plastic hinge pins on either side of the display housing. Release them by turning the slotted head anti-clockwise with a small flat-bladed screwdriver, then pull them straight out.



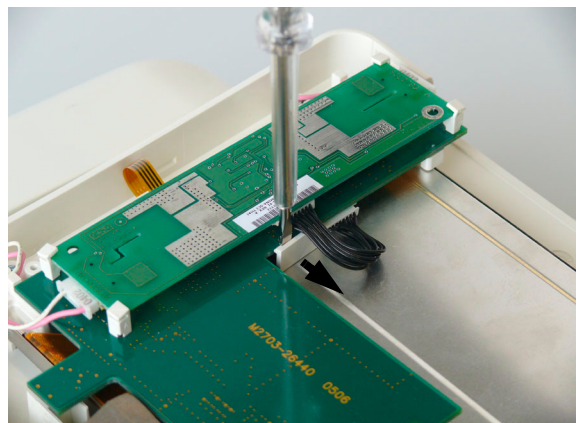
- 3 Starting at the hinge end of the display assembly, separate the two halves of the display assembly.



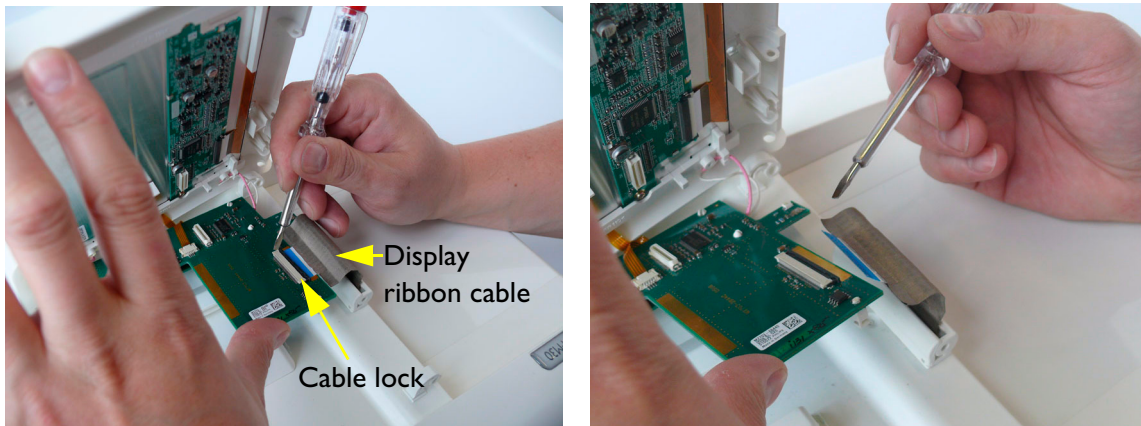
- 4 Open up the two halves of the display assembly and then remove the back of the display unit.



- 5 Disconnect the backlight inverter cable after gently releasing it with the help of a small, flat-bladed screwdriver.

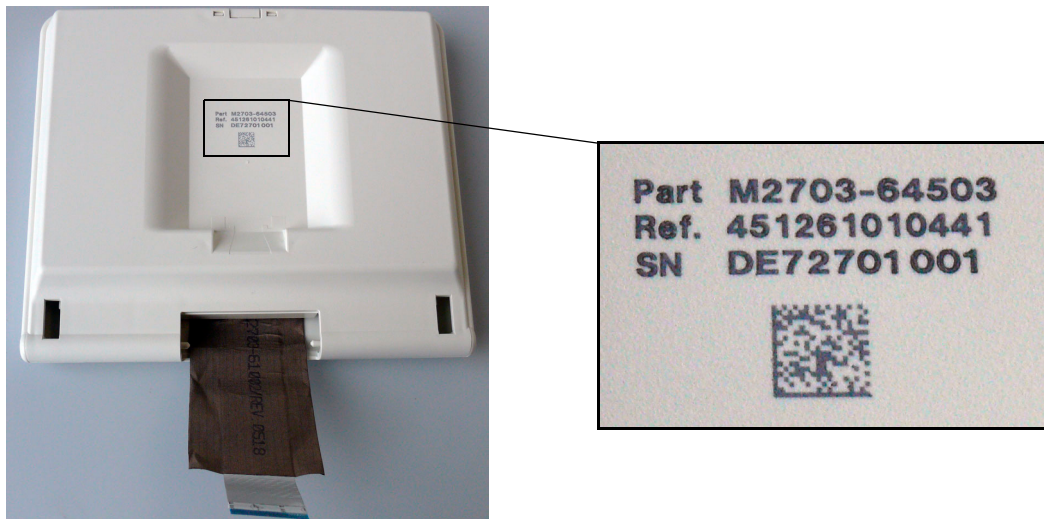


- 6 Disengage the cable lock for the display ribbon cable on the display assembly side using a small, flat-bladed screwdriver, then disconnect the ribbon cable. You will reuse this original ribbon cable to connect the new display assembly (after discarding the ribbon cable that comes with the new display assembly). You have now removed the display assembly.



## Replacing the Display Assembly

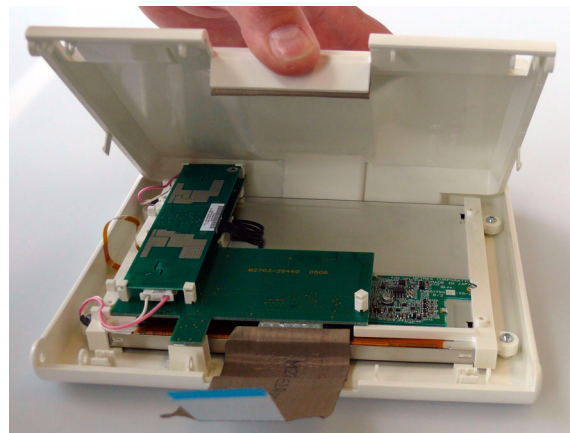
The new display assembly has the part number, reference (ordering) number, and the serial number printed on the rear.



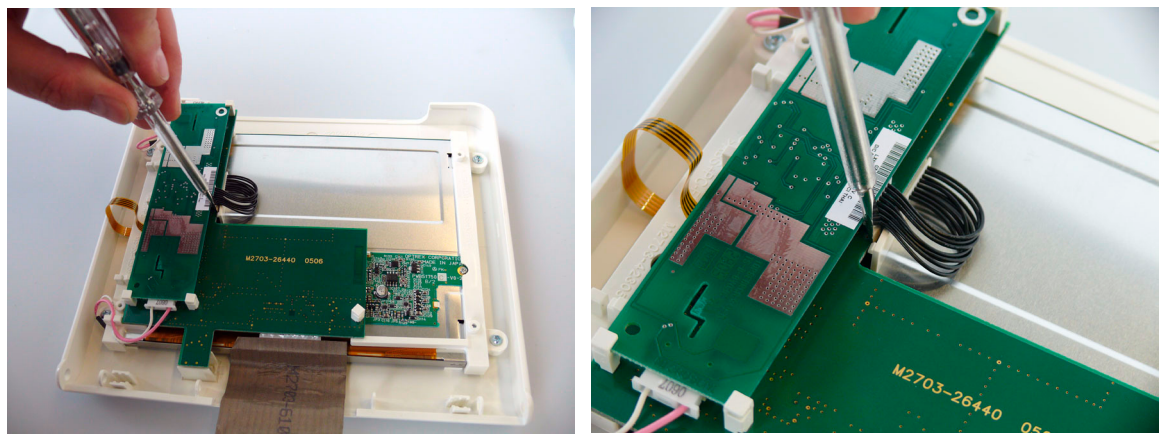
- 1 Remove the two plastic hinge pins on either side of the new display housing. Release them by turning the slotted head anti-clockwise with a small flat-bladed screwdriver, then pull them straight out.



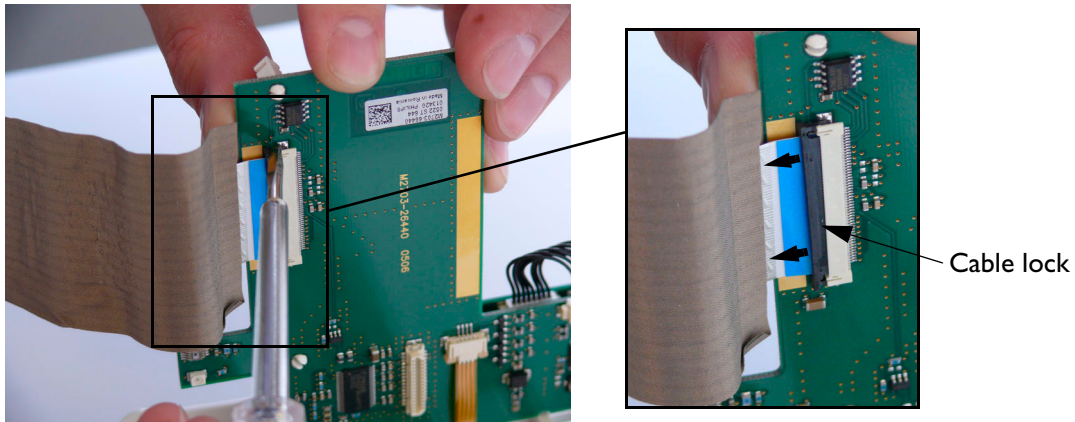
- 2 Open up the new display assembly, and remove the rear of the display housing.



- 3 Disconnect the backlight inverter cable after gently releasing it with the help of a small, flat-bladed screwdriver.



- 4 Disengage the cable lock for the display assembly ribbon cable with the help of a small, flat-bladed screwdriver. Remove the cable, and discard it. (You will reuse the original ribbon cable, still attached to the monitor, for connecting the new display assembly.)



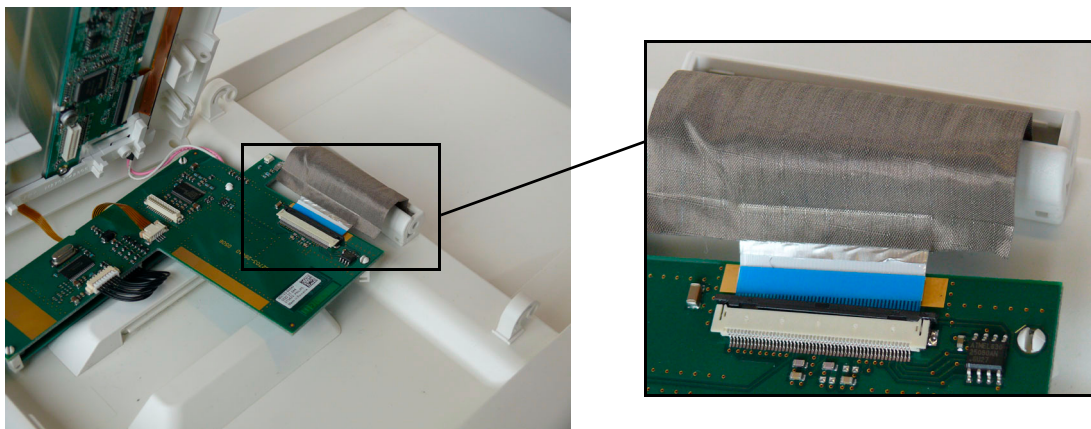
- 5 Remove the cable, and discard it. (You will reuse the original ribbon cable, still attached to the monitor, for connecting the new display assembly.)



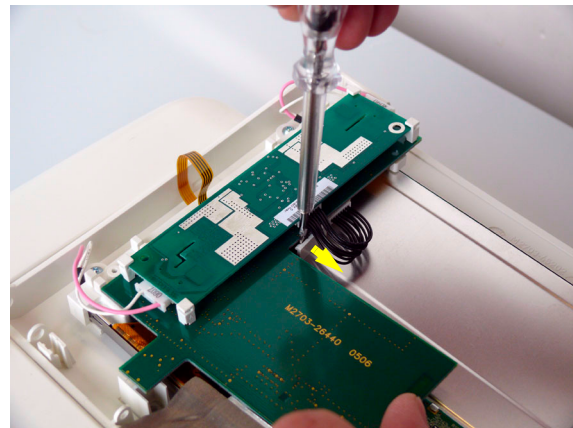
Discard ribbon cable from new display assembly and recycle according to local laws



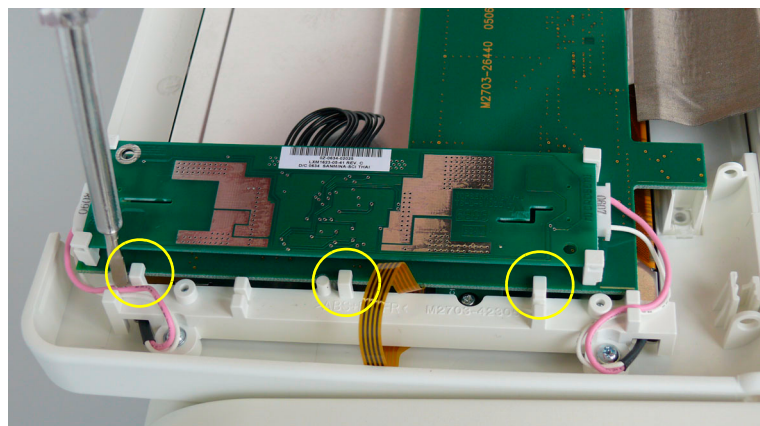
- 6 Connect the open end of the original display ribbon cable (attached to the monitor) to the connector on the new display assembly. Ensure the cable lock is engaged and the cable is firmly connected.



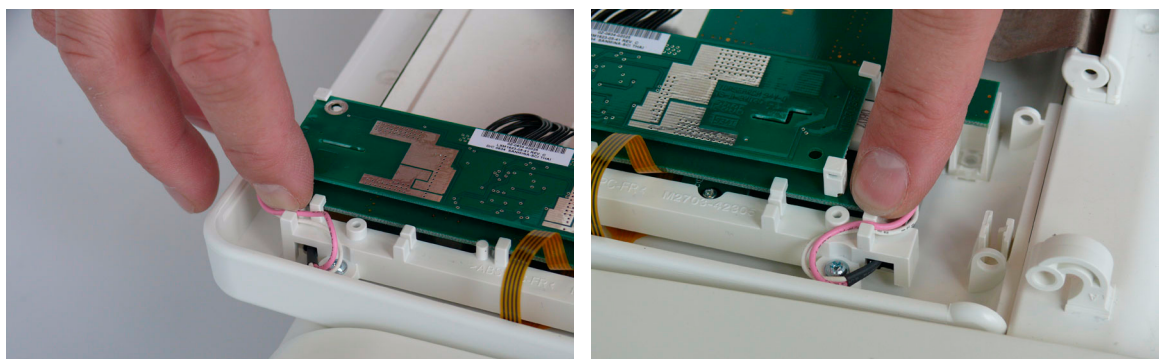
- 7 Reconnect the backlight converter cable, and slot the PCBs back into position. Use a small, flat-bladed screwdriver to very gently lever move the backlight converter connector outwards (in the direction of the arrow) to guide the PCB back into position. Otherwise the bPCB would catch on the connector.



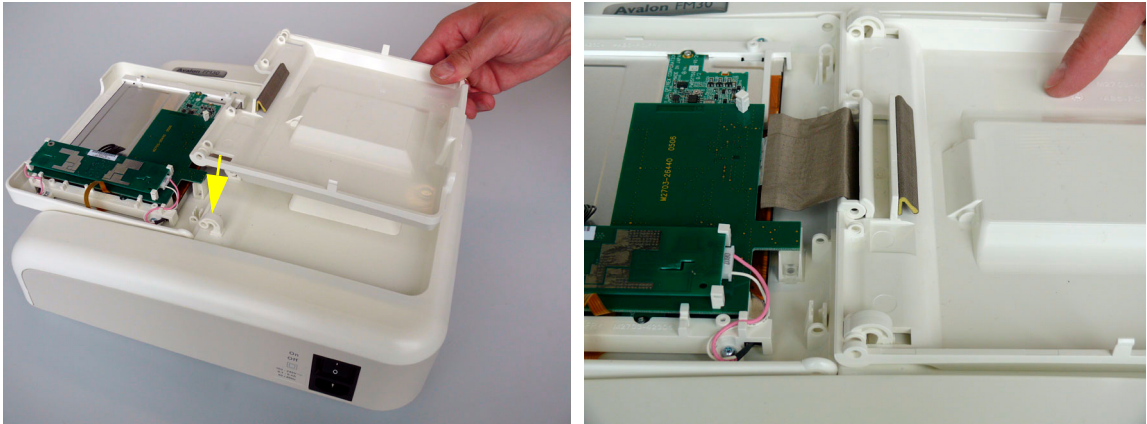
- 8 Ensure the PCB is located correctly as shown in the photograph...



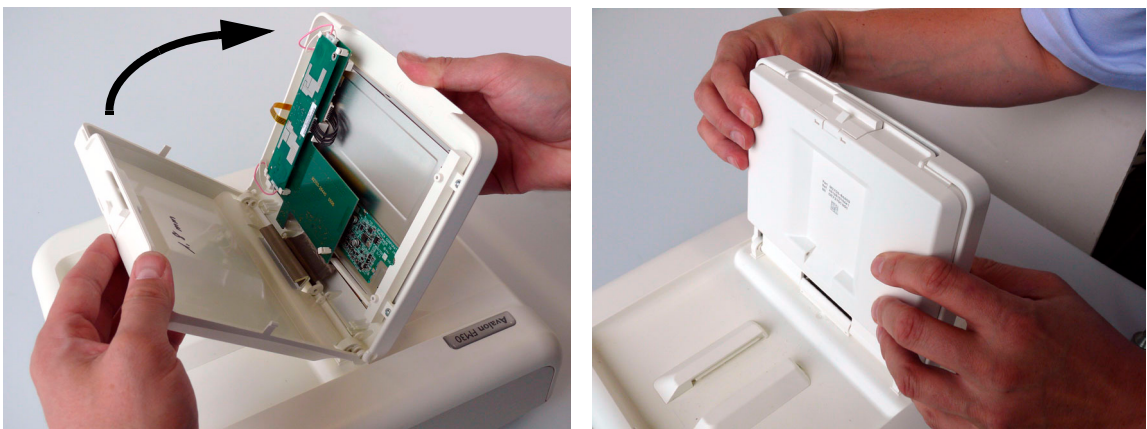
...and that the pink and white cables are correctly routed, and held in place.



9 Take the new rear housing cover and place it in the position shown.



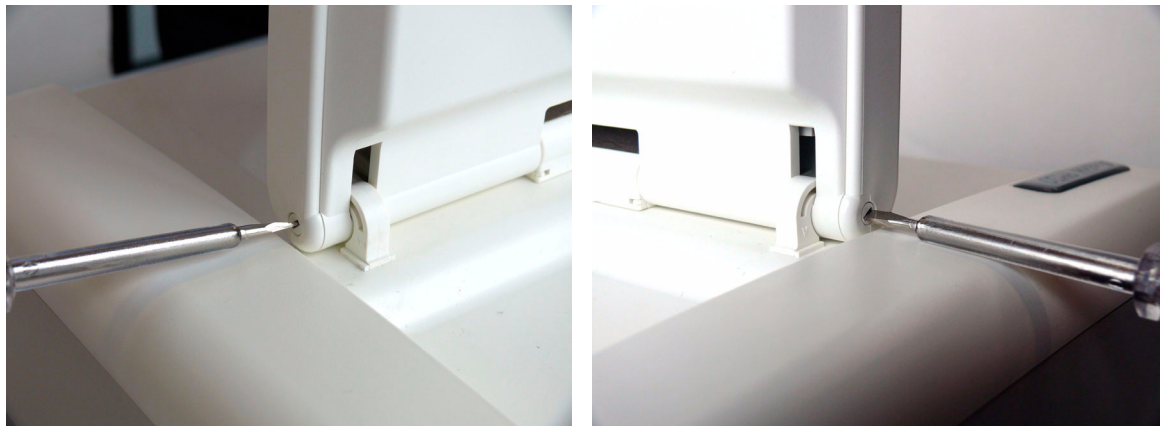
10 Put the two halves of the display assembly together as shown.



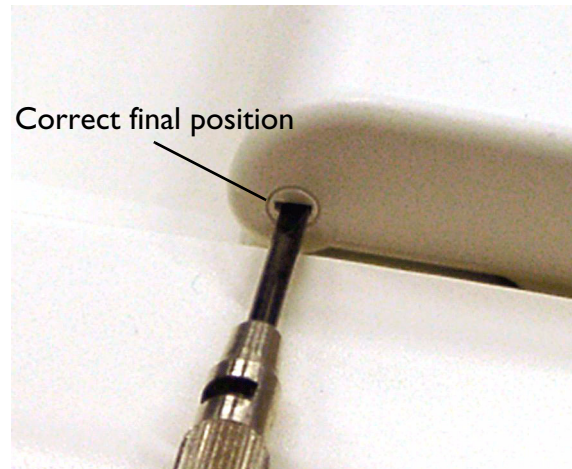
11 Insert the new plastic hinge pins.



12 Do not apply excessive force when refitting the hinge pins. Turn the head of the hinge pin gently with a small screwdriver. You will feel when the pin is seated correctly, and the head should be flush with the surface.



The slot in the head of the pin should be pointing in the same plane as the side of the display assembly housing when the display is folded flat.



13 Reattach the ratchet clip to the top cover of the monitor and to the rear of the display housing.



## Recorder Disassembly

The recorder consists of the following major sub-assemblies:

- Drawer Assembly
- Recorder Chassis
- Thermal Line Printhead (TLPH) Holder
- Recorder Adapter Board
- Stepper Motor

### Removing the Drawer Assembly

- 1 Press the paper table release to unlock the paper drawer and then pull the table forward to open it fully.

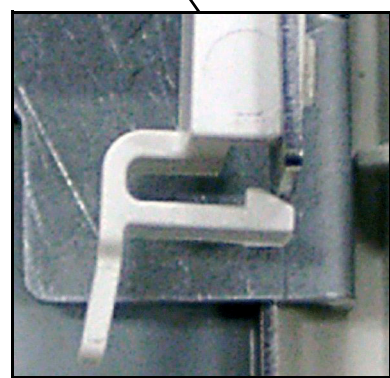
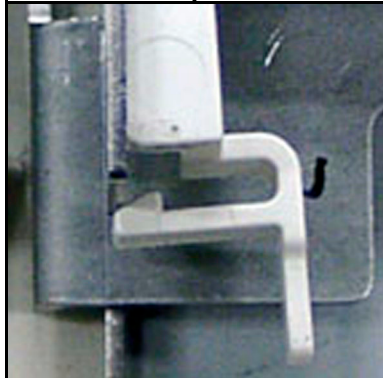


- 2 Squeeze the two plastic lugs on the underside of the drawer to release the drawer, and then pull to remove the drawer. (Here shown with the top cover removed.)



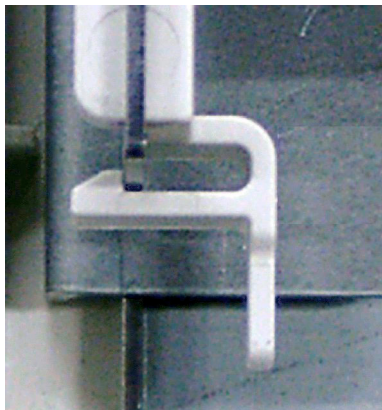


Detail showing plastic hooks open

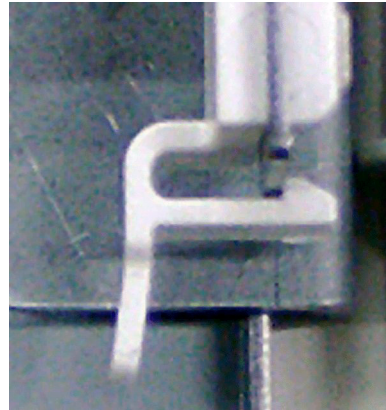


## Replacing the Drawer Assembly

Before replacing the drawer assembly, refer to the previous photographs, and study the detail showing the position of the plastic hooks when the drawer is fixed in position.

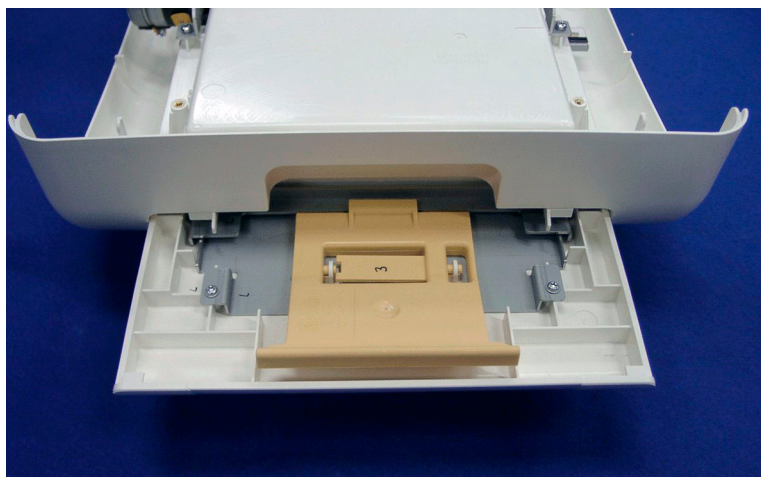
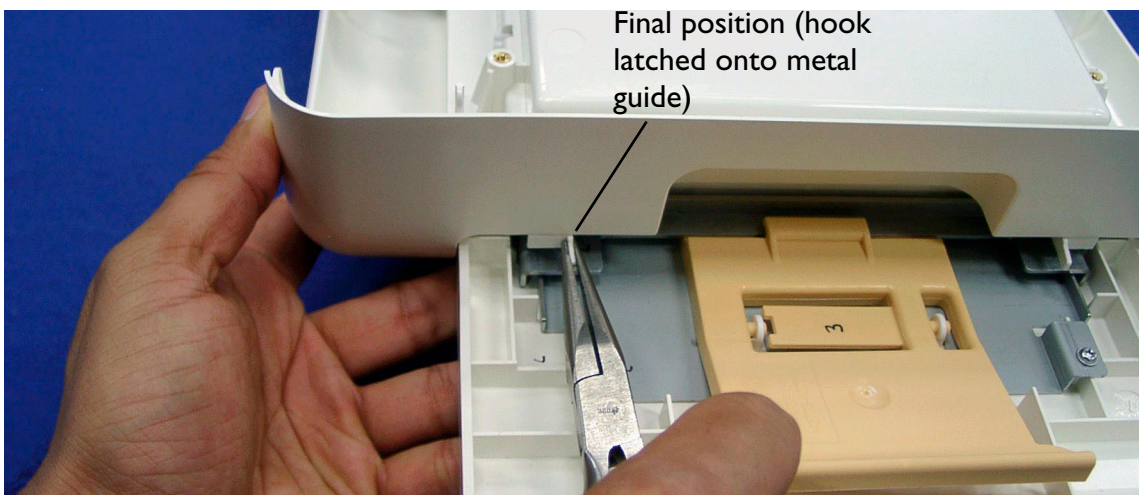


Detail showing plastic hooks latched onto metal guide. This is how the hooks should be when the drawer is fixed in position.



- 1 Slide the drawer into the drawer recess on the top cover, and check that it is located correctly on the runners.
- 2 Latch the two plastic hooks onto the metal guides to secure the drawer in place. You will find it easier to use a long-nosed pair of pliers to move the hook into position, as shown in the following photographs.



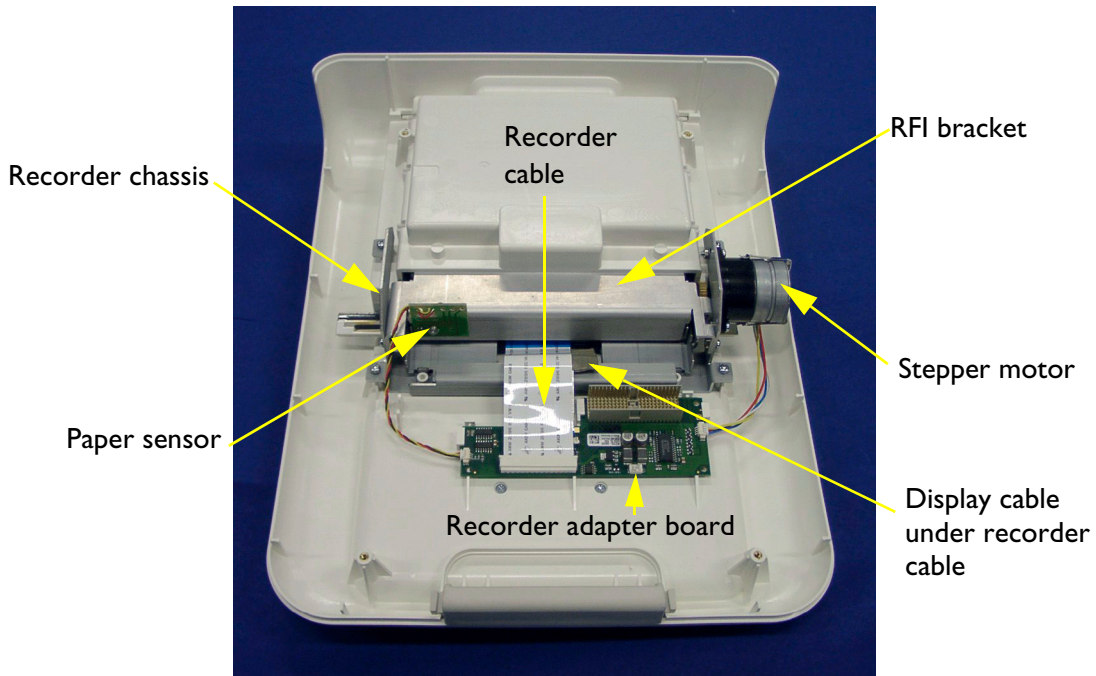


## Removing the Recorder Chassis

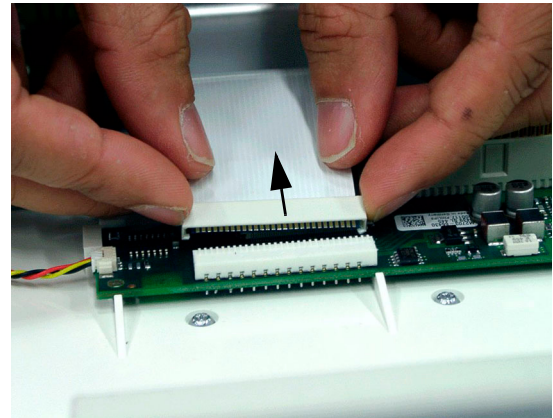
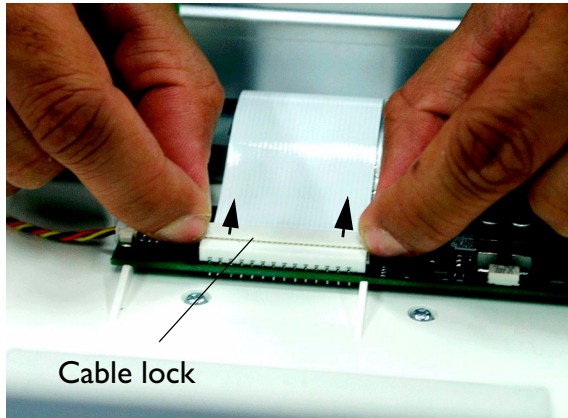
- 1 Remove the top cover assembly (see page 105).
- 2 Slide open the paper drawer, to gain access to the two countersunk screws (in the following photographs, we have removed the drawer assembly).
- 3 Remove the two countersunk screws.



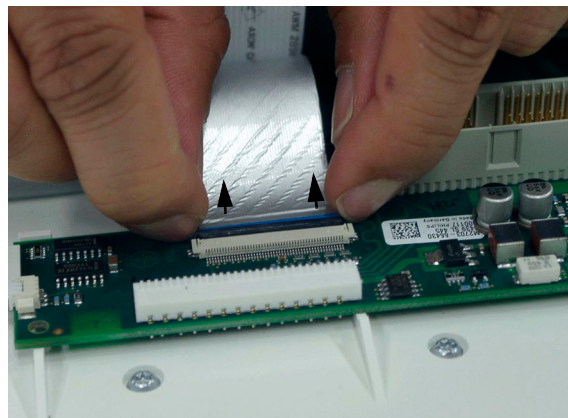
- 4 Turn over the top cover assembly and place it top down on a cloth or other soft surface.



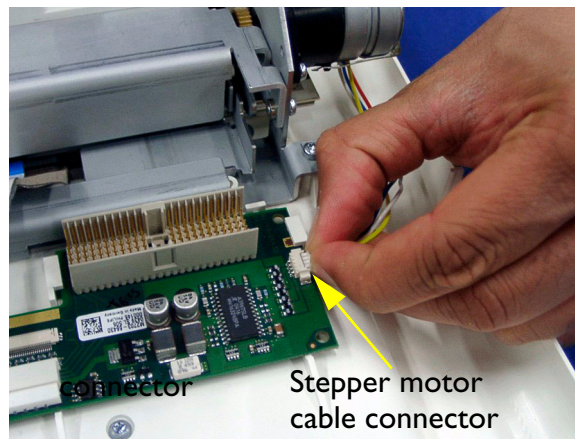
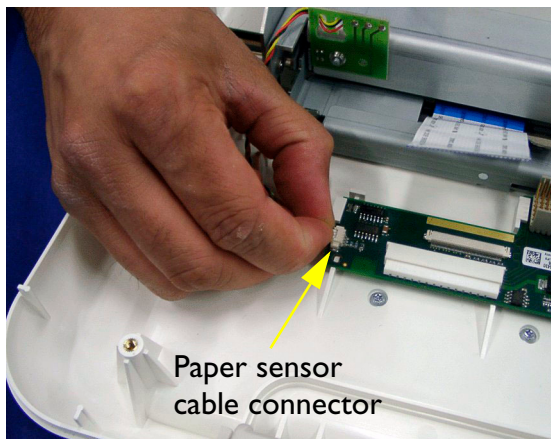
- 5 First release the cable lock by moving it in the direction of the arrows, then disconnect the white recorder ribbon cable from the recorder adapter board.



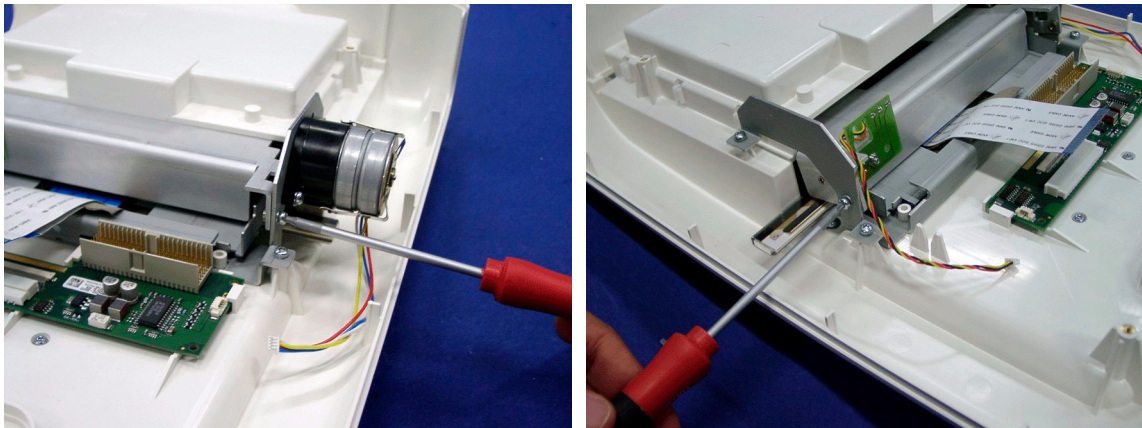
- 6 Next, disconnect the silver display ribbon cable from the recorder adapter board after releasing the cable lock.



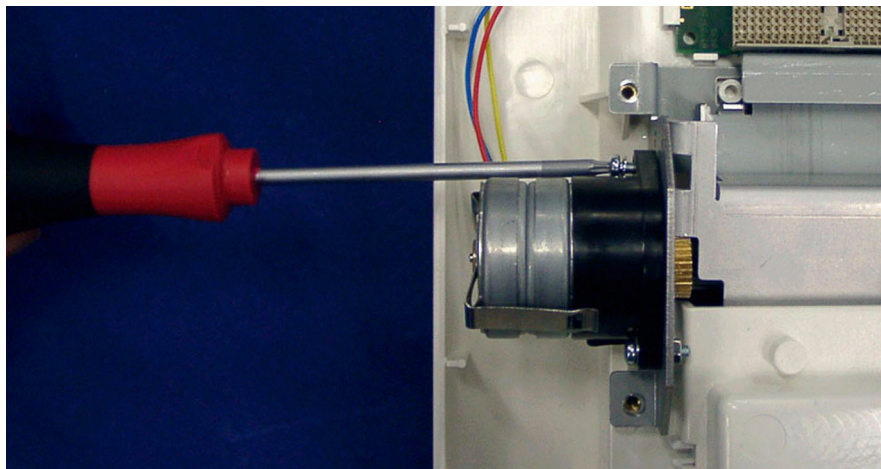
- 7 Disconnect the stepper motor the paper sensor cable connectors from the recorder adapter board.



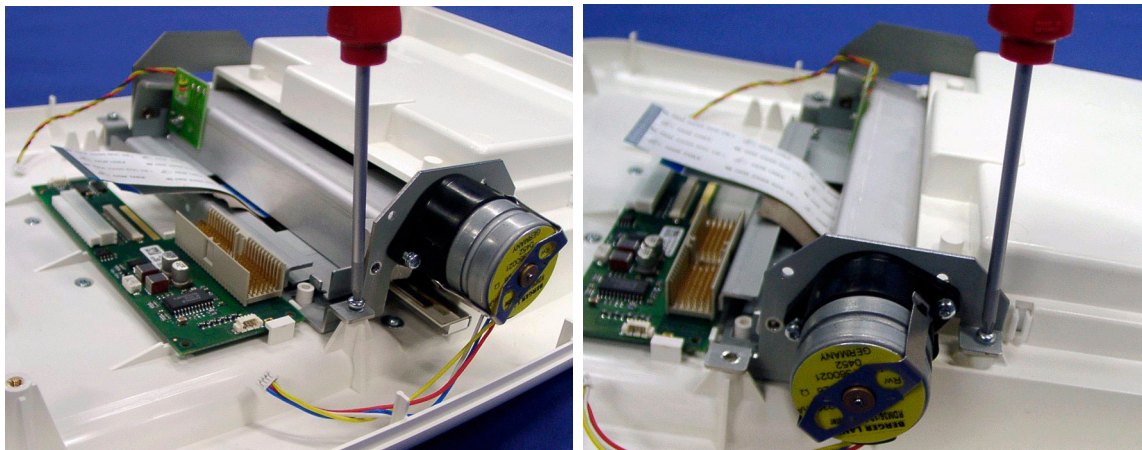
- 8 Remove the two screws (one on each side) fastening the RFI bracket to the recorder chassis.

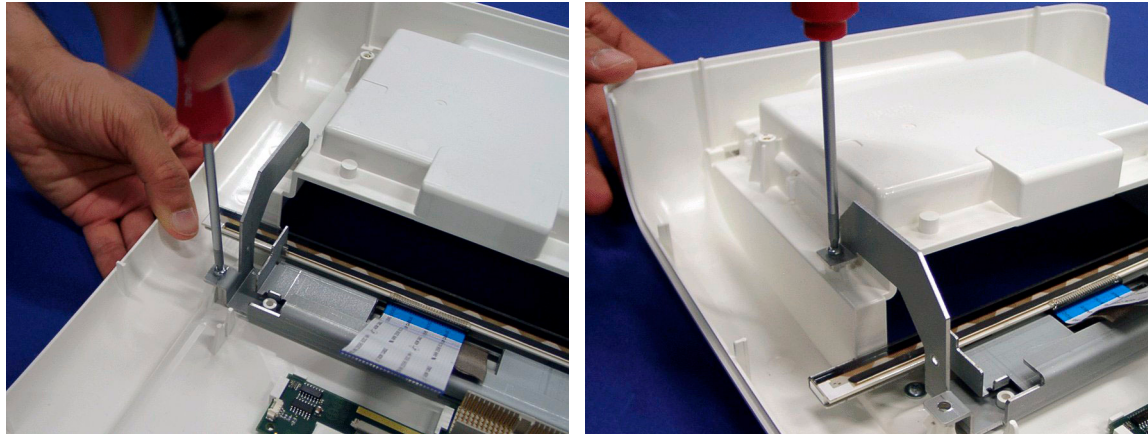


- 9 Partially unscrew the screw nearest the recorder adapter board holding the stepper motor to the recorder chassis, until the thread no longer protrudes. This allows the necessary clearance for removing the RFI bracket.

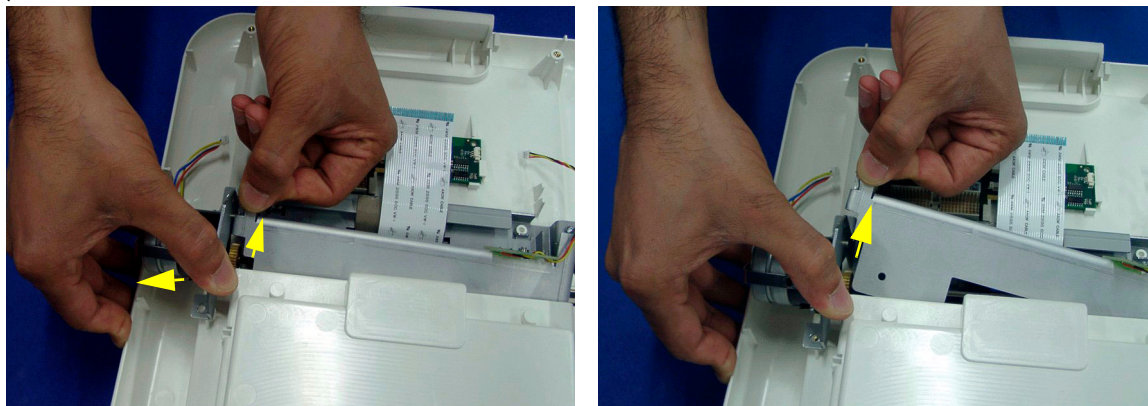


- 10 Remove the four screws holding the recorder chassis.

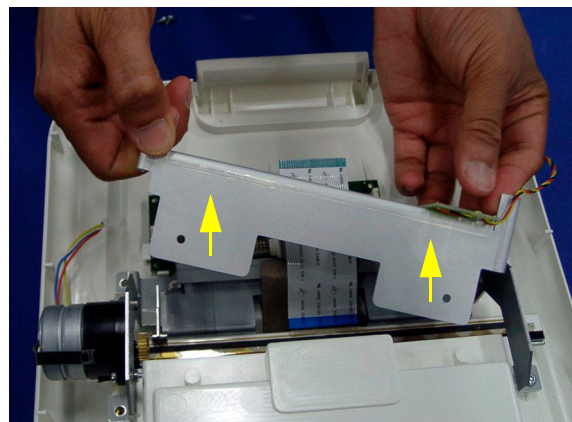




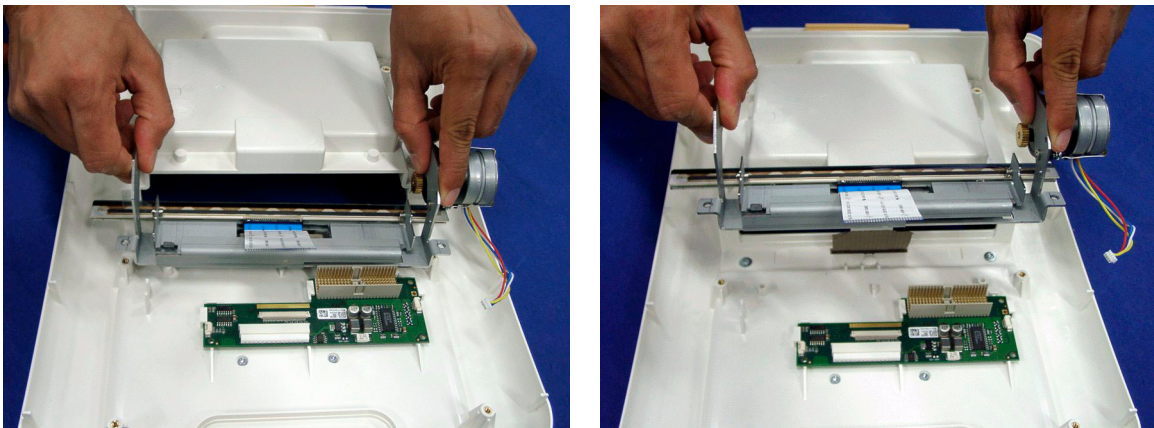
11 Free the RFI bracket on the stepper motor side by applying a little sideways pressure to the recorder chassis, while at the same time pulling the RFI bracket forwards.



12 Remove the RFI bracket.

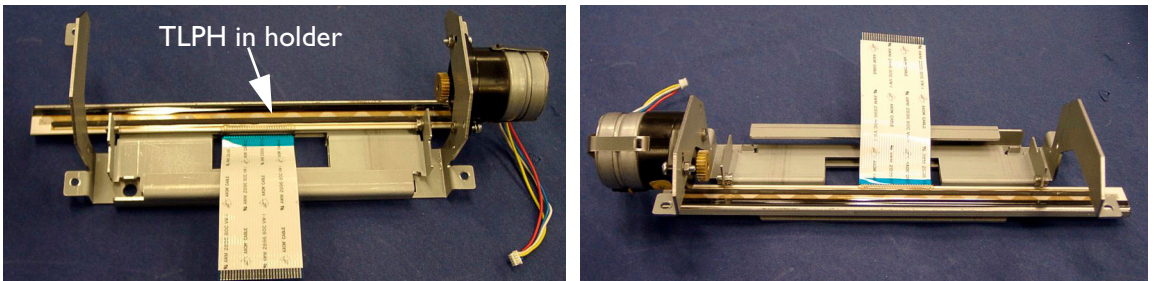


13 Lift out the recorder chassis.

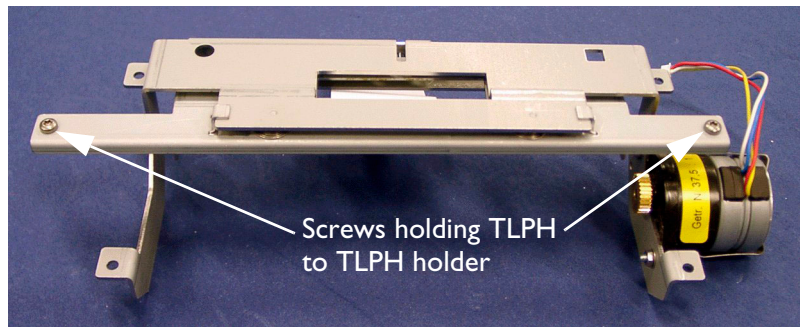


### Removing the Thermal Line Printhead (TLPH)

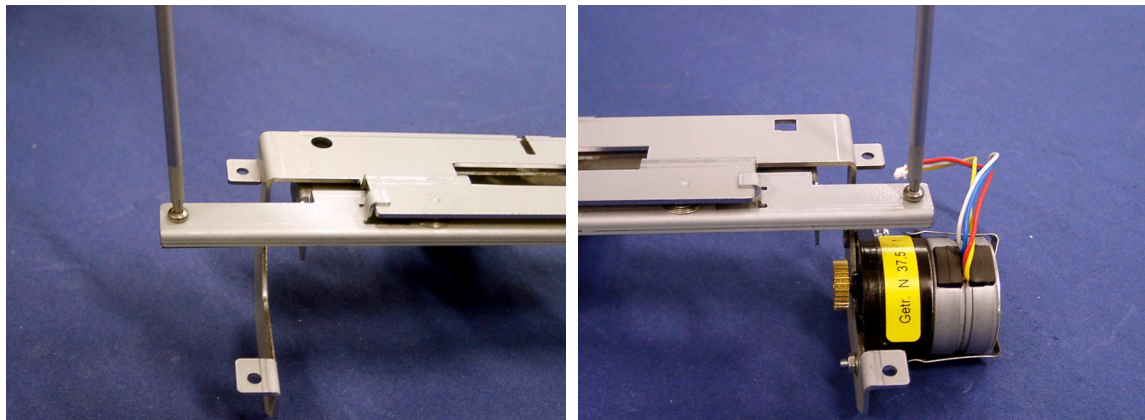
1 Remove the recorder chassis as described in the section “Replacing the Recorder Chassis” on page 126.



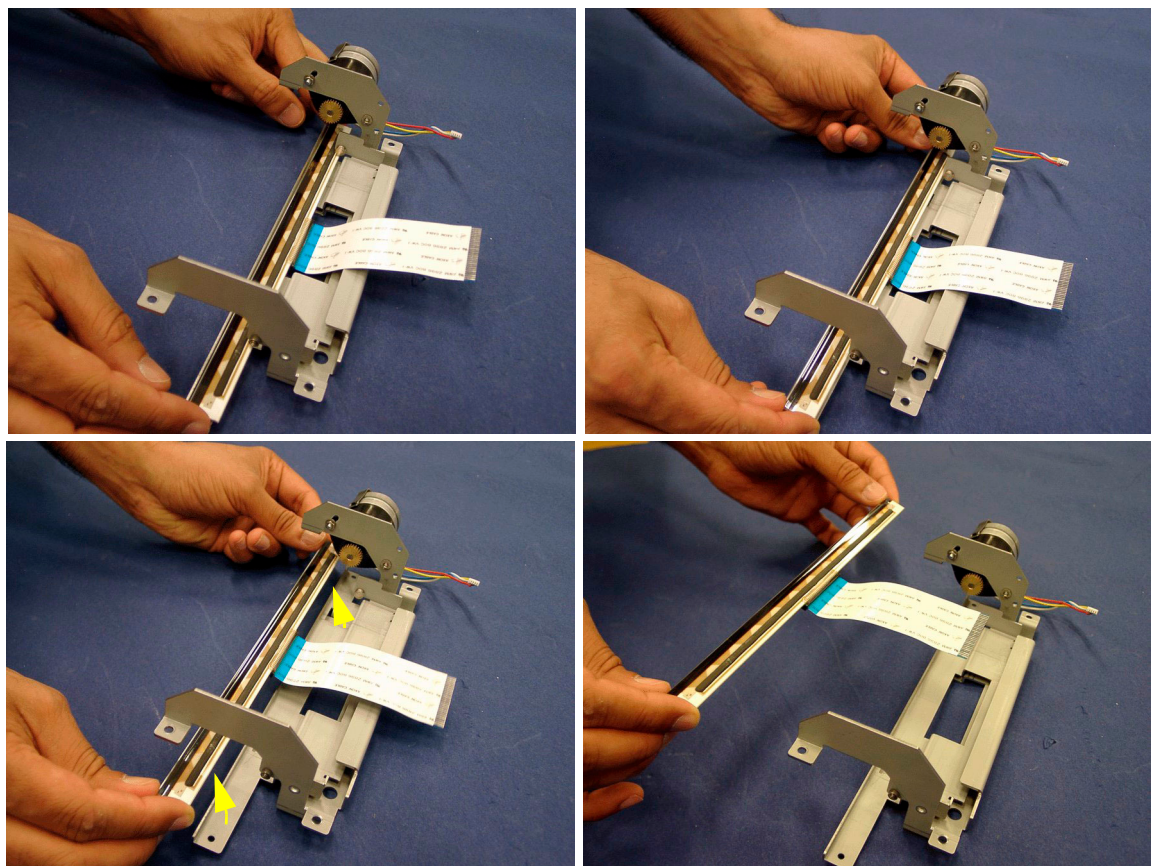
2 Turn the recorder chassis over.



- 3 Remove the two screws holding the TLPH to the TLPH holder.



- 4 Turn the chassis assembly over again, and carefully remove the TLPH from the holder as shown.



### Replacing the TLPH

The procedure for replacing the TLPH is a reversal of the removal procedure.

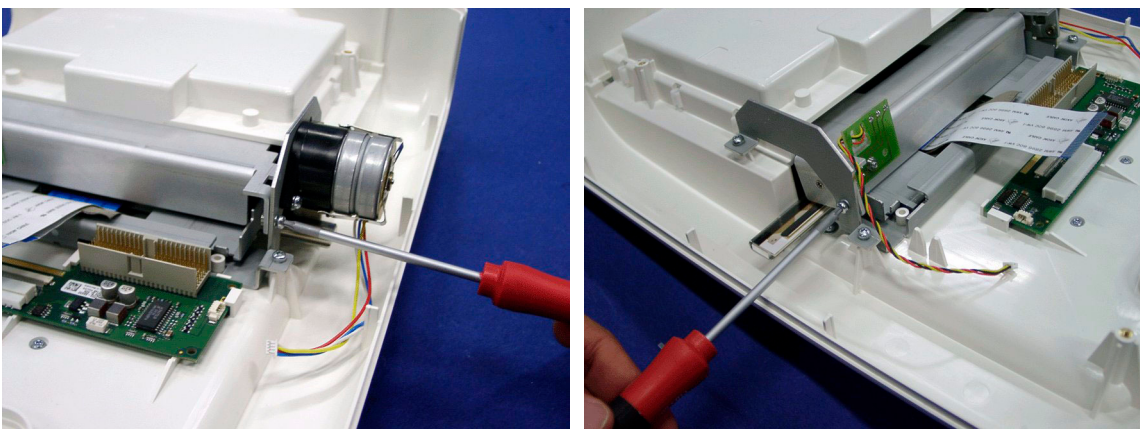
## Replacing the Recorder Chassis

The procedure for replacing the recorder chassis is a reversal of the removal procedure (see “Replacing the Recorder Chassis” on page 126 for the sequence).

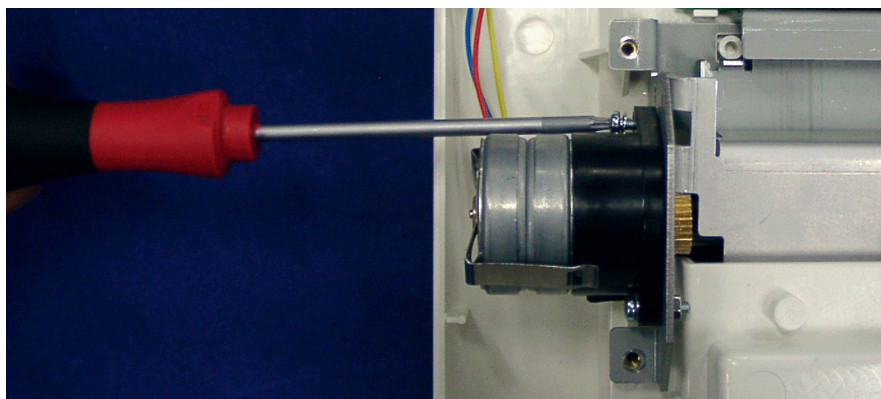
- 1 Ensure that the TLPH holder has been fitted to the recorder chassis, as described in the previous section.
- 2 Place the recorder chassis into position, and drive the four fixing screws in lightly. Do not tighten yet!
- 3 Turn over the top cover housing, then secure the two countersunk screws. This centers the recorder chassis correctly.



- 4 Turn over the top cover again, and replace the RFI bracket, reversing the removal procedure. Secure the RFI bracket to the recorder chassis with the two screws.



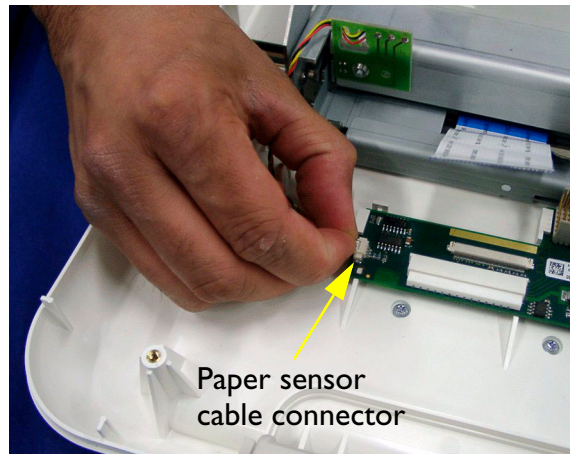
- 5 Tighten the stepper motor screw that you loosened to allow removal of the RFI bracket.



- 6 Now tighten the four screws to secure the recorder chassis.
- 7 Ensure that you reconnect the paper sensor cable and the stepper motor cable to the recorder adapter board.

## Removing the Paper Sensor Assembly

- 1 Remove the top cover assembly (see “Top Cover Disassembly/Reassembly” on page 105).
- 2 Place the top cover assembly top down on a cloth or other soft surface.
- 3 Disconnect the paper sense cable connector from the recorder adapter board.



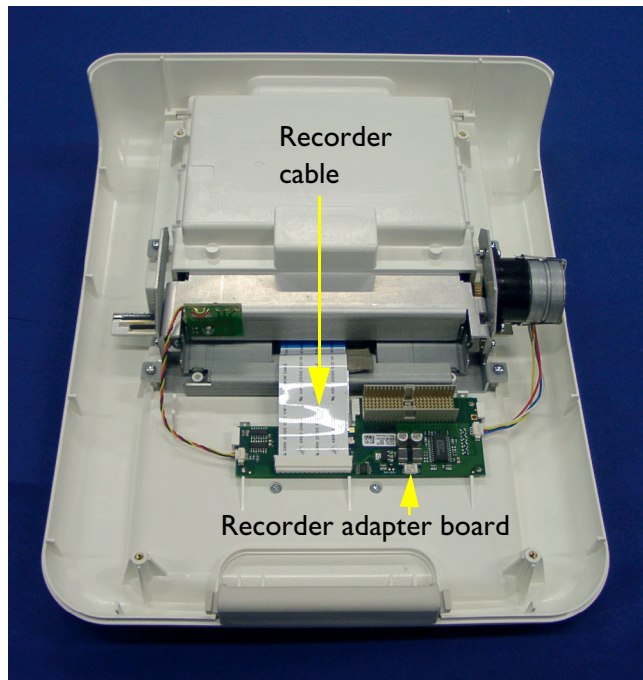
- 4 Turn over the top cover and remove the screw holding the paper sensor to the RFI bracket.

## Replacing the Paper Sensor Assembly

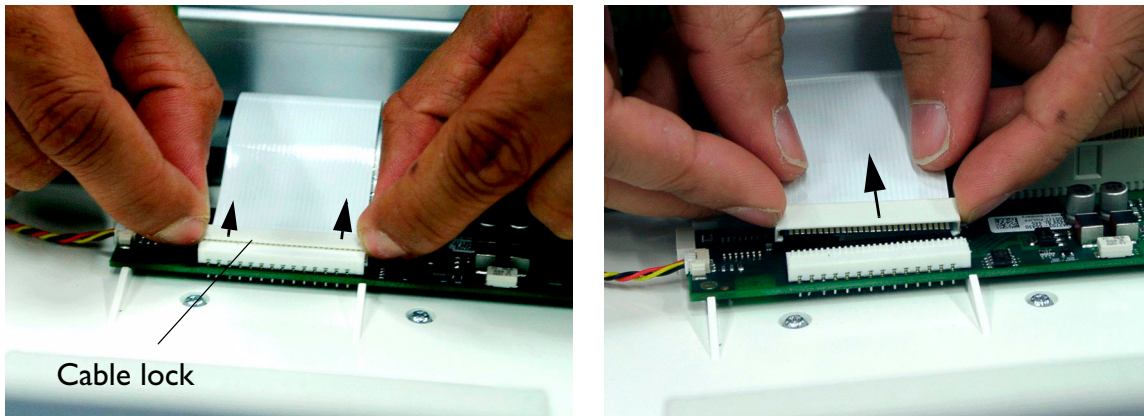
The procedure for replacing the paper sensor is a reversal of the removal procedure. Ensure that the paper sensor cable is properly connected to the recorder adapter board.

## Removing the Recorder Adapter Board

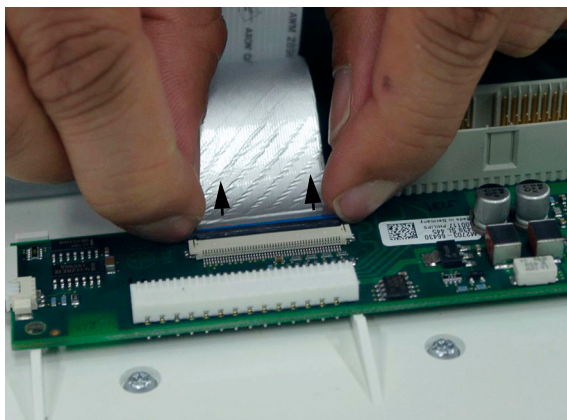
- 1 Remove the top cover assembly (see “Top Cover Disassembly/Reassembly” on page 105).
- 2 Place the top cover assembly top down on a cloth or other soft surface.



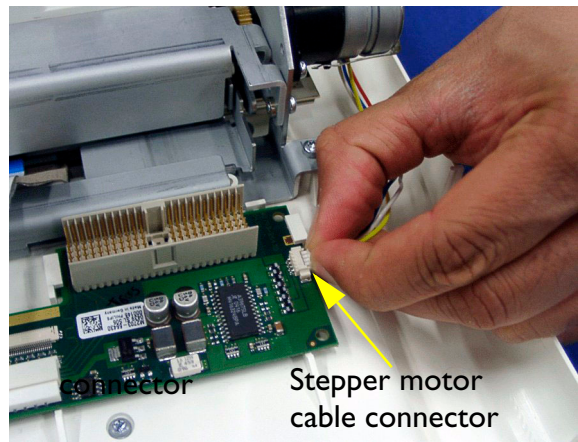
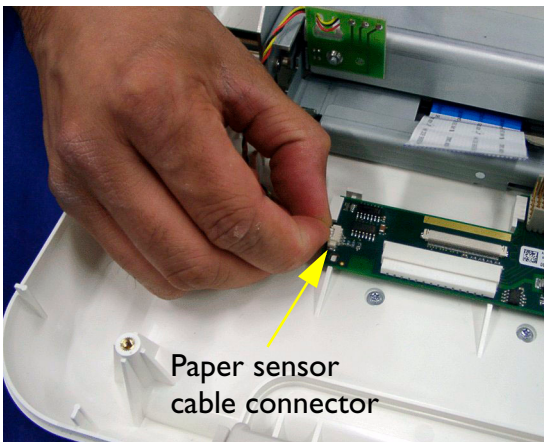
- 3 First release the cable lock by moving it in the direction of the arrows, then disconnect the white recorder ribbon cable from the recorder adapter board.



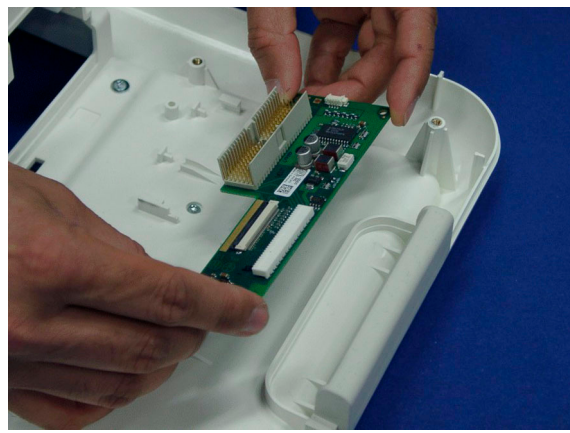
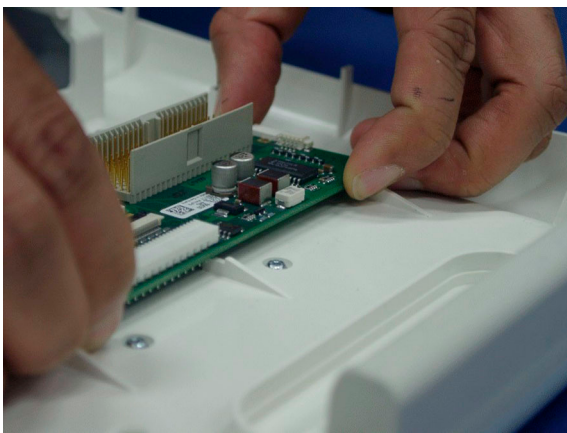
- 4 Next, disconnect the silver display ribbon cable from the recorder adapter board after releasing the cable lock.



- 5 Disconnect the stepper motor the record-sense cable connectors from the recorder adapter board.



- 6 With all cables disconnected, remove the recorder adapter board.

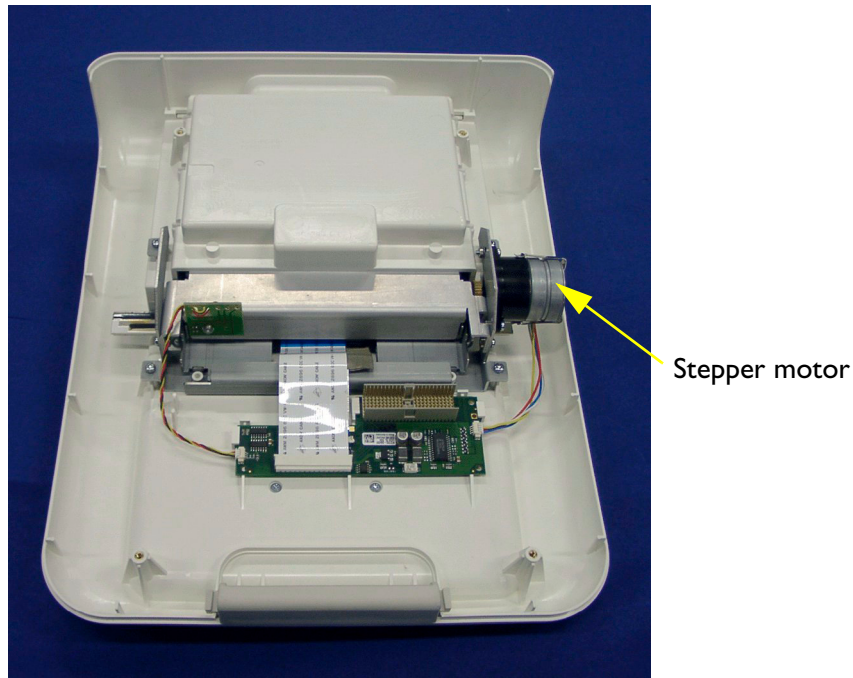


## Replacing the Recorder Adapter Board

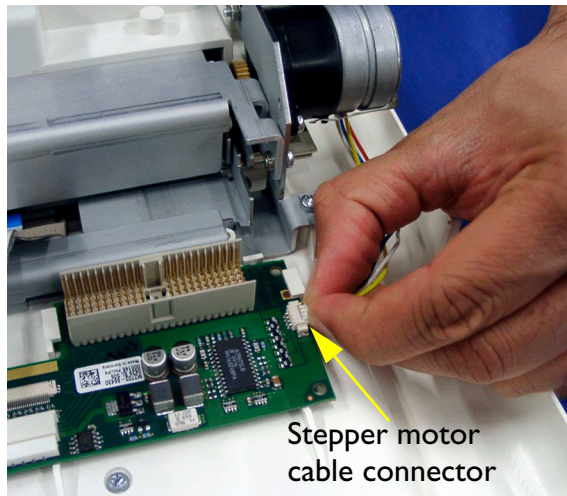
The procedure to replace the recorder adapter board is a reversal of the removal procedure. Ensure that all cables are firmly reconnected.

## Removing the Stepper Motor

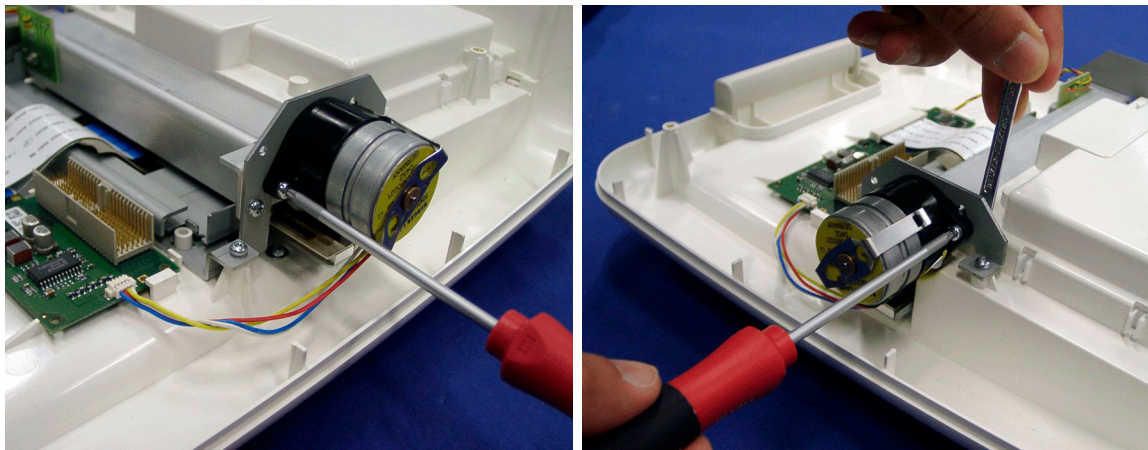
- 1 Remove the top cover assembly (see “Top Cover Disassembly/Reassembly” on page 105).
- 2 Place the top cover assembly top down on a cloth or other soft surface.



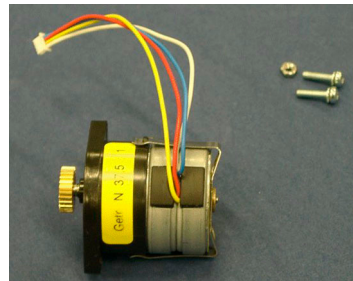
- 3 Disconnect the stepper motor cable connector from the recorder adapter board.



- 4 Remove the two screws holding the stepper motor to the recorder chassis. Note that the upper screw is secured with a small nut.



- 5 Remove the stepper motor.



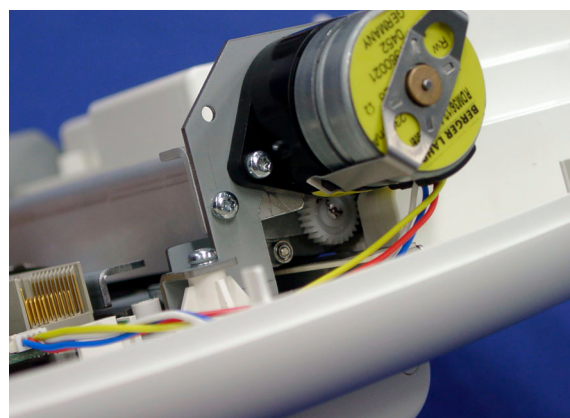
## Replacing the Stepper Motor

- 1 Refit the stepper motor to the recorder chassis, but do NOT tighten the screws yet! The upper slot on the stepper motor allows fine adjustment when meshing the gears together.
- 2 Close the drawer assembly.
- 3 Gently mesh the stepper motor gear with that of the paper roller.

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**CAUTION** Do NOT press the gears together or exert any pressure on the stepper motor spindle.

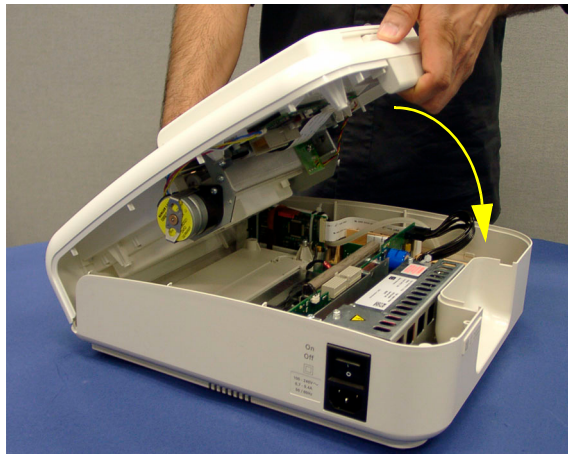
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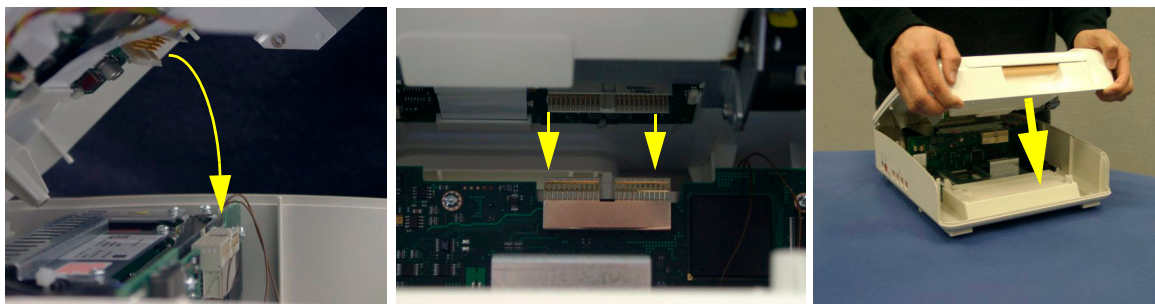
- 4 When in place, tighten the screws, remembering that the upper screw is secured with a small nut.

## Replacing the Top Cover Assembly

- 1 Ensure all items are replaced in the top cover assembly. Check that all the cables are reconnected.
- 2 Carefully align the front edge of the top cover assembly with the front edge of the bottom housing assembly.



- 3 With the front edge of the top cover still located in the front edge of the bottom housing, gently lower the top cover, making sure the multi-pin connector on the recorder adapter board aligns with the socket on the main CPU board.



- 4 Place the top cover back to its normal position. Apply a little pressure to seat the multi-pin connector.

- 5 Holding both assemblies together, carefully place the unit upside down on a soft surface to prevent scratching or other damage.



- 6 Refit the four screws securing the top cover assembly to the bottom housing, using a T-10 Torx driver, as a reversal of the procedure in “Top Cover Disassembly/Reassembly” on page 105. Turn the monitor the right way up.

**CAUTION** When replacing the top cover, do not over-torque the screws. Excessive torque may damage the screw mountings.

## Bottom Cover Disassembly/Reassembly

This section describes disassembly and reassembly operations for the bottom cover and its assemblies.

### Removing the Power Supply Assembly

- 1 Remove the top cover assembly (see page 105).
- 2 Disconnect the power supply cable connector from the main CPU board. While removing the connector, support the end of the main CPU board to prevent excessive flexing.

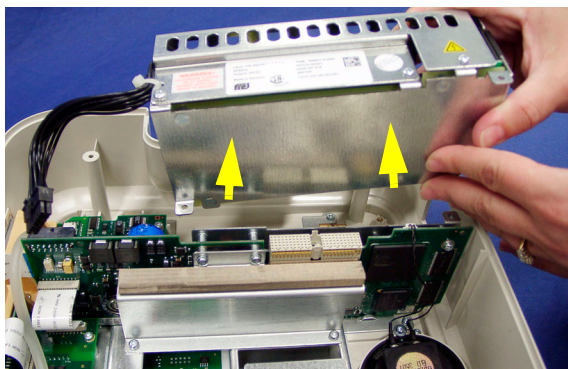
Remove power supply cable



- 3 Remove the three screws securing the power supply



- 4 Lift the cable end of the power supply assembly with one hand, while guiding the power socket/on/off switch free of the aperture in the bottom housing, then lift out the power supply

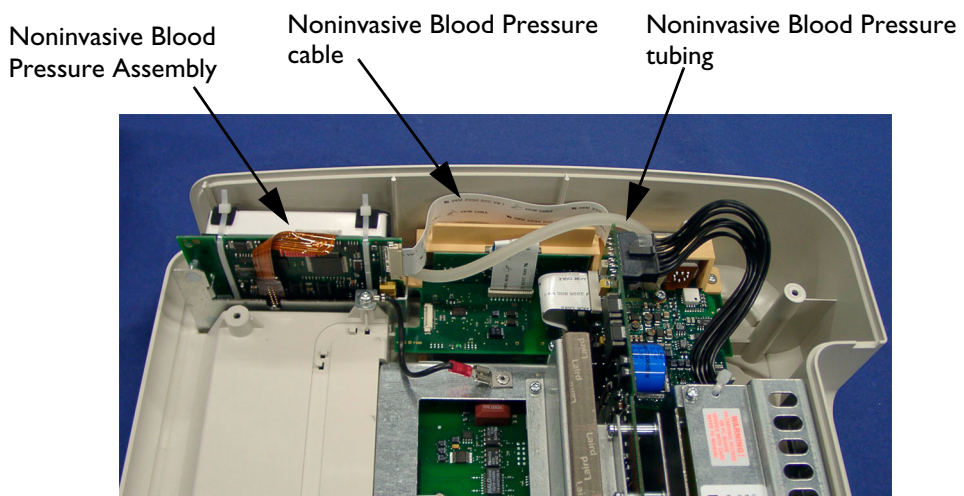


## Replacing the Power Supply Assembly

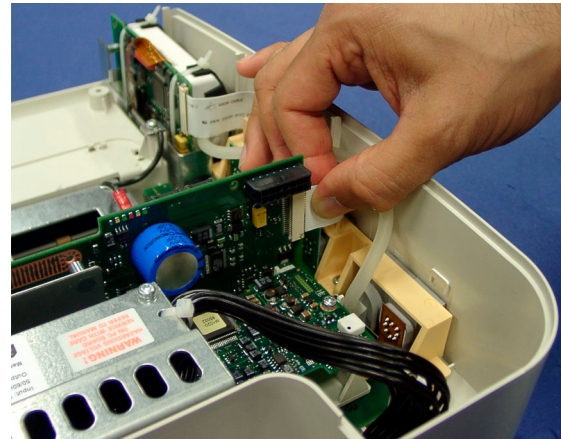
The procedure to replace the power supply assembly is a reversal of the removal procedure. Remember to reconnect the power supply cable connector.

## Removing the Noninvasive Blood Pressure Assembly

- 1 Remove the top cover assembly (see page 105).  
The NiBP assembly is identified in the next picture.



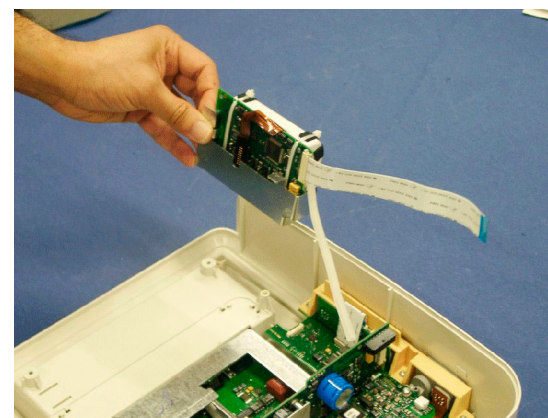
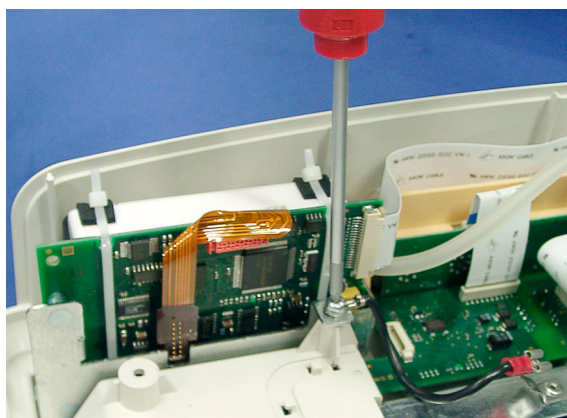
- 2 Disconnect the tubing from the noninvasive blood pressure connector, and the ribbon cable from the main CPU board (after first disengaging the cable lock).



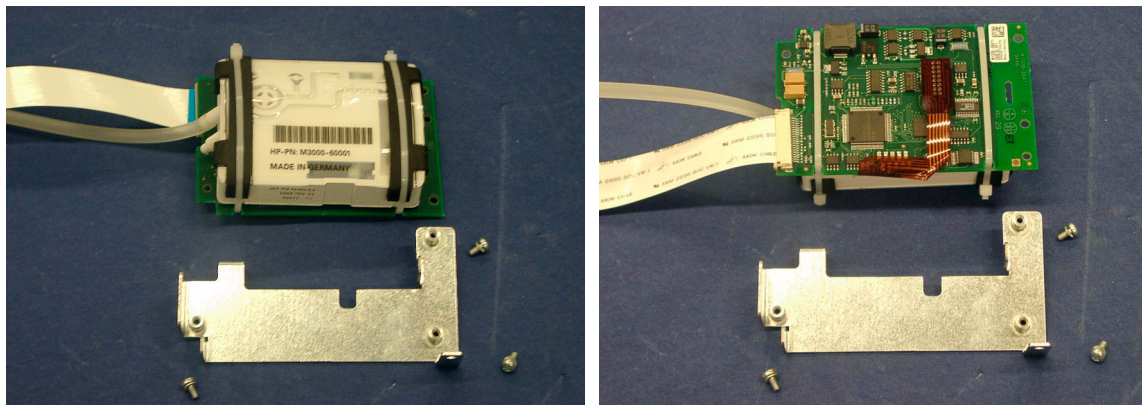
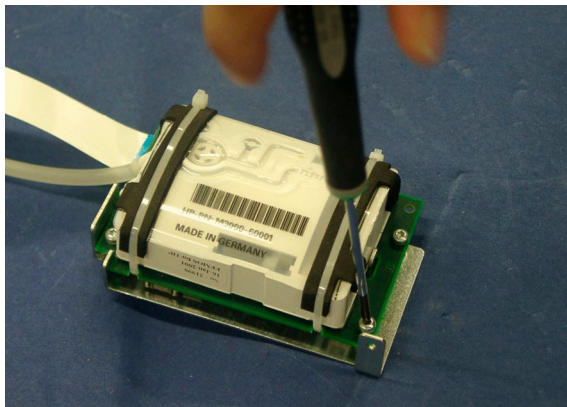
- 3 Disconnect the grounding cable's spade connector.



- 4 Remove the two screws holding the noninvasive blood pressure assembly to the bottom housing, then remove the noninvasive blood pressure assembly.

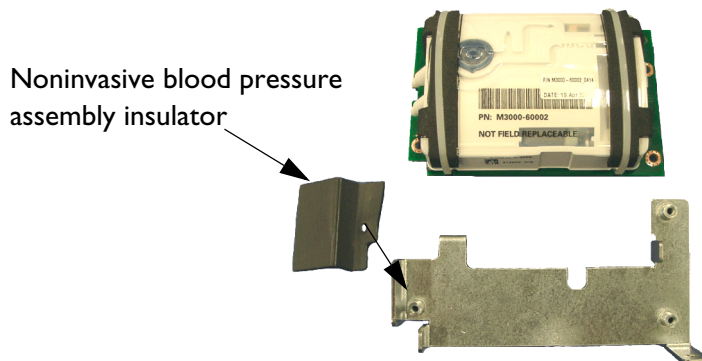


- 5 To separate the noninvasive blood pressure assembly from the noninvasive blood pressure assembly holder, remove the three screws.



## Replacing the Noninvasive Blood Pressure Assembly

- 1 The new noninvasive blood pressure assembly kit comes with an insulator. Place the insulator on the metal holder as shown.



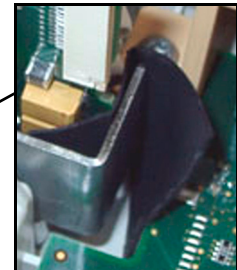
- 2 Place the noninvasive blood pressure assembly onto the metal holder, ensuring the insulator is in place.



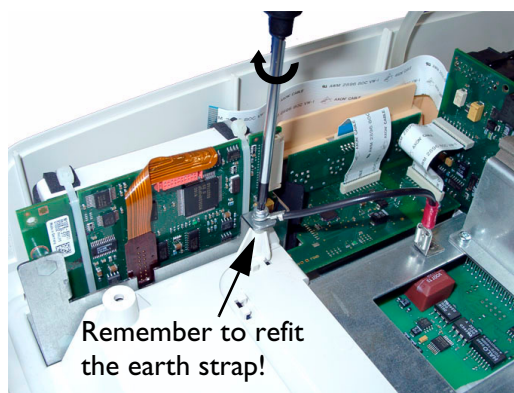
- 3 Refit the three screws securing the NiBP module to the holder.



- 4 Place the noninvasive blood pressure assembly into position, making sure that the insulator is located correctly as shown.



- 5 Refit the earth strap.

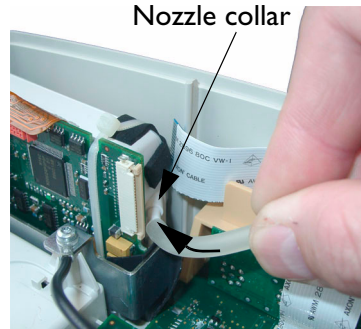


- 6 Refit the remaining screw to secure the noninvasive blood pressure assembly to the bottom housing.

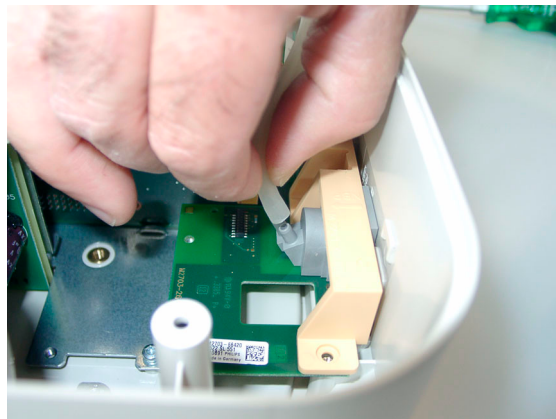


- 7 Refit the noninvasive blood pressure tubing first (before reconnecting the ribbon cable).

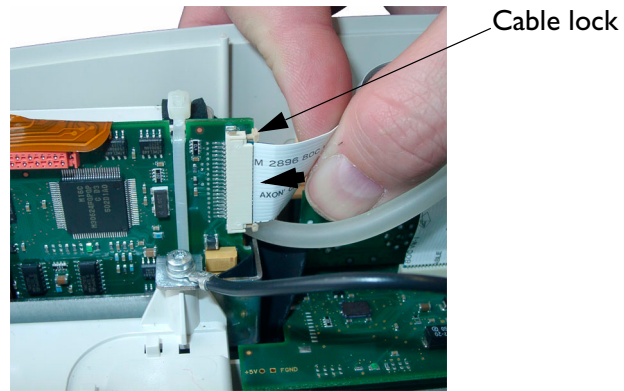
Reconnect the noninvasive blood pressure tubing first. Push it right up to the nozzle collar, and check that it is secure.



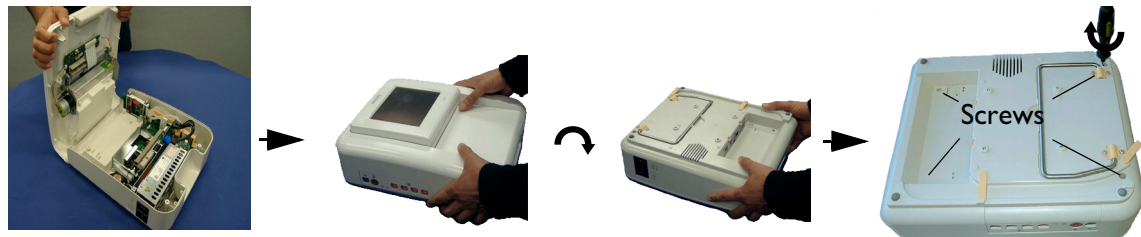
When refitting the tubing, also make sure you connect it to the left-hand connector when facing the front of the sockets.



- 8 Then reconnect the ribbon cable. Make sure the cable lock is firmly closed and the cable is secure.

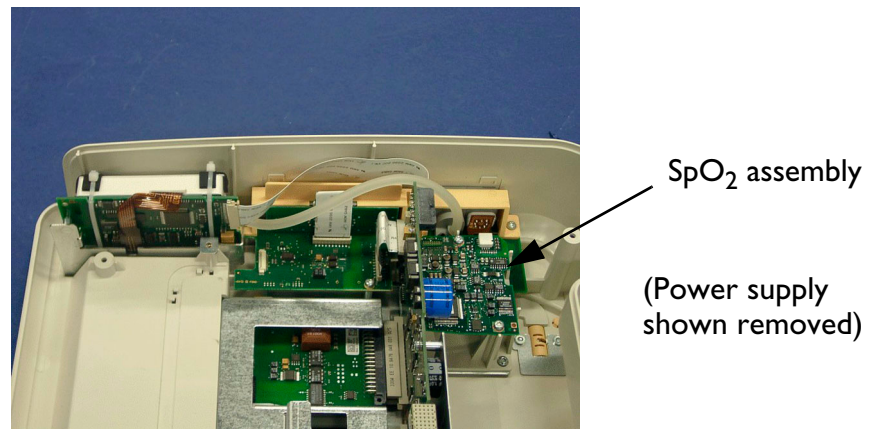


- 9 Replace the top cover (see page 132).

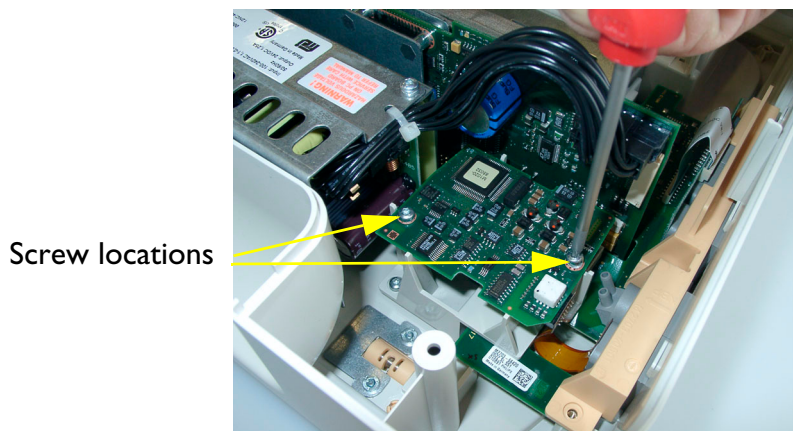


## Removing the SpO<sub>2</sub> Assembly

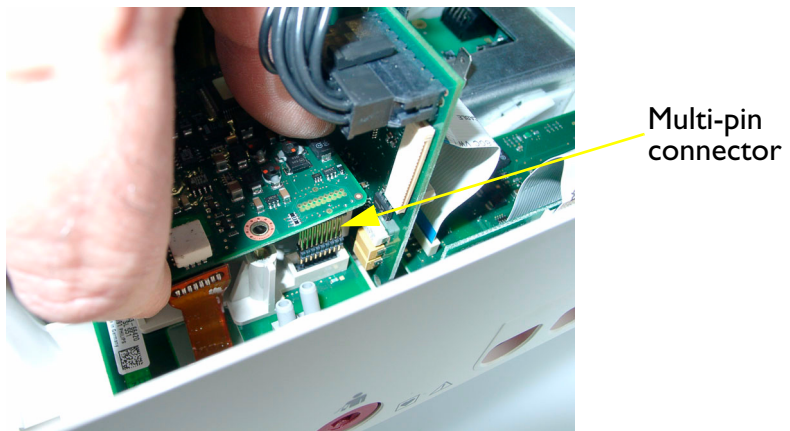
- 1 Remove the top cover assembly (see page 105).  
The SpO<sub>2</sub> assembly is identified in the next picture.



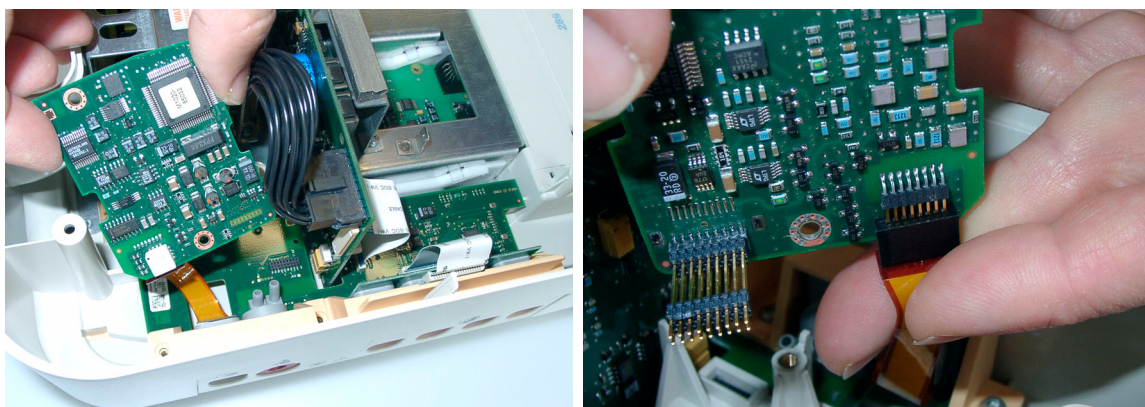
- 2 Remove the two screws holding the SpO<sub>2</sub> assembly.



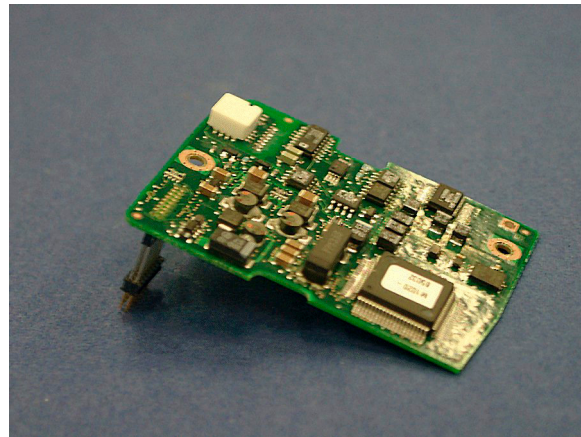
- 3 Lift the side of the SpO<sub>2</sub> assembly nearest the SpO<sub>2</sub> socket, carefully disconnecting the multi-pin connector shown.



- 4 To remove the flat brown SpO<sub>2</sub> socket cable from the multi-pin connector on the underside of the SpO<sub>2</sub> board, it is easier to lift the board first to get better access.



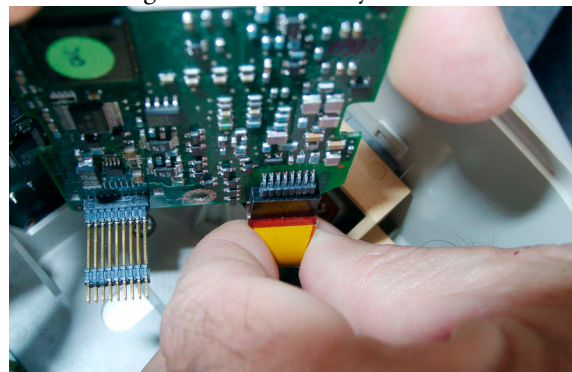
- 5 Remove the SpO<sub>2</sub> assembly.



## Replacing the SpO<sub>2</sub> Assembly

The procedure to replace the SpO<sub>2</sub> assembly is a reversal of the removal procedure.

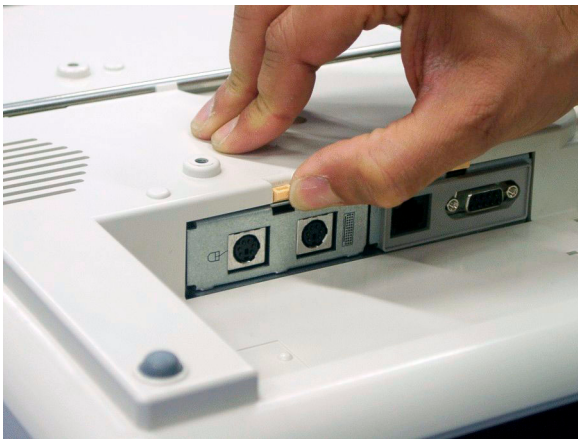
Hint: When reconnecting the flat brown SpO<sub>2</sub> socket cable to the multi-pin connector on the SpO<sub>2</sub> board, it is easiest to reconnect it while holding the board vertically in line with the cable connector as shown.



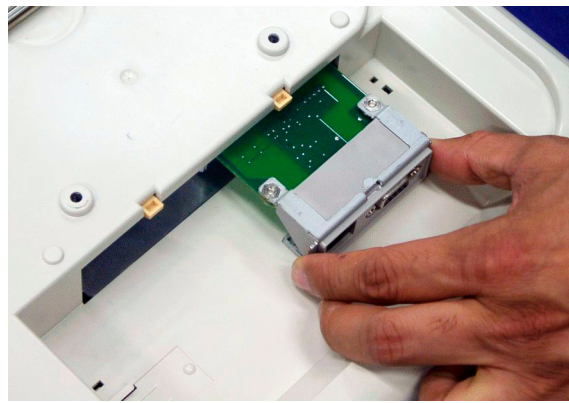
## Removing the Interface Boards

The interface boards can be accessed from the underside of the monitor's housing.

- 1 Turn the monitor upside down on a non-scratch surface.
- 2 Release the board by pressing the clip that keeps the board in place, as shown. Use a small flat-headed screwdriver to gently prise the board out if it is too tight to pull out with your fingers.



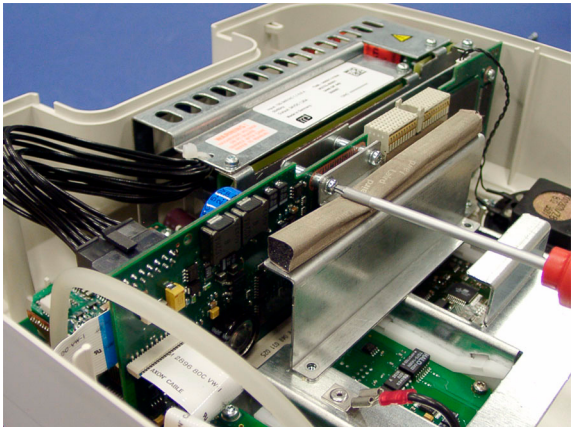
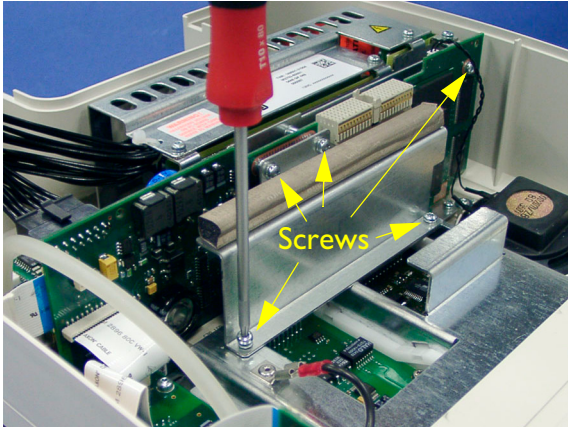
3 Pull the board out.



## Removing the Main CPU Board

To remove the main CPU board, proceed as follows:

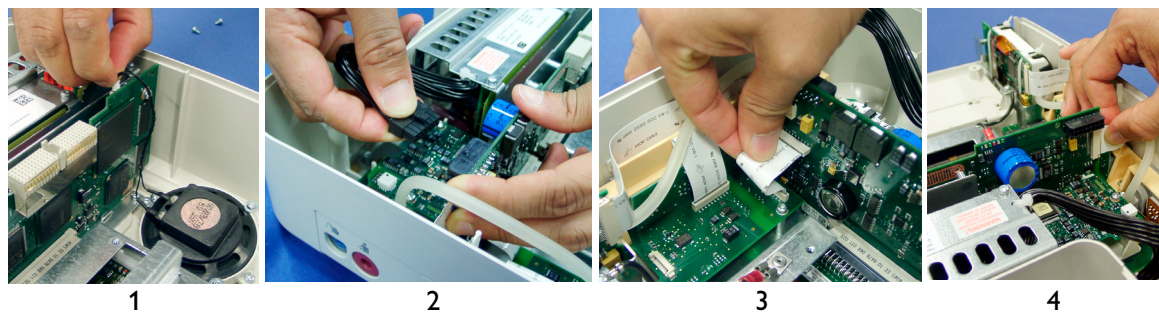
- 1 Remove the top cover assembly (see page 105).
- 2 Remove the screws holding the metal shield.



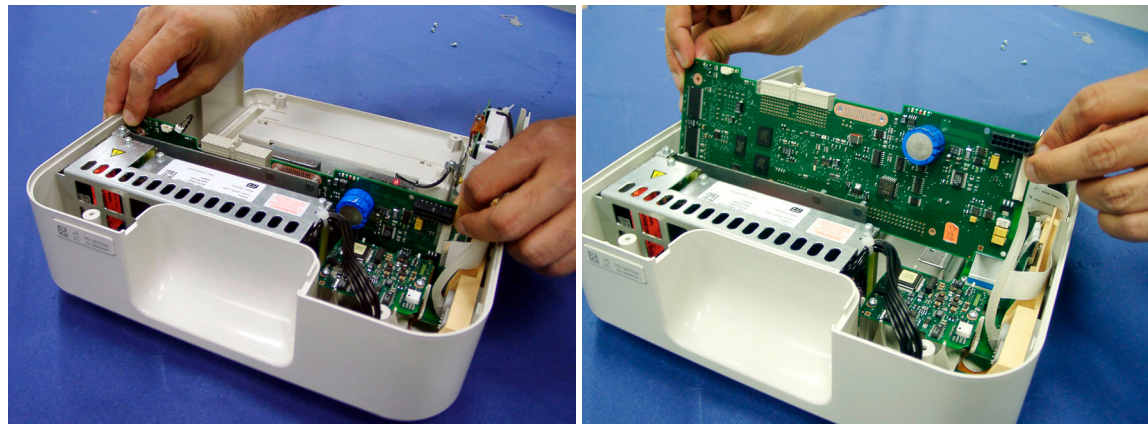
- 3 Remove the shield.



- 4 Disconnect the loudspeaker cable (1), the power supply cable (2), the bus master board cable (3), and the noninvasive blood pressure cable (4) from the main CPU board.



- 5 Remove the main CPU board by lifting it straight up.



## Replacing the Main CPU Board

The procedure to replace the main CPU board is a reversal of the removal procedure. Ensure all the cables are properly reconnected.

## Removing the Front End Assembly

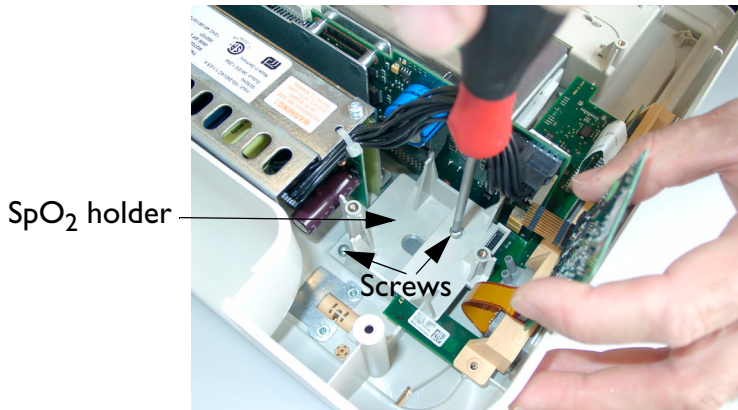
The front end assembly consists of:

- the fetal sensor socket connectors with ribbon cable
- the bus master board with ribbon cable

- the optional noninvasive blood pressure and SpO<sub>2</sub> socket connectors
- the connector block frame

To remove front end assembly:

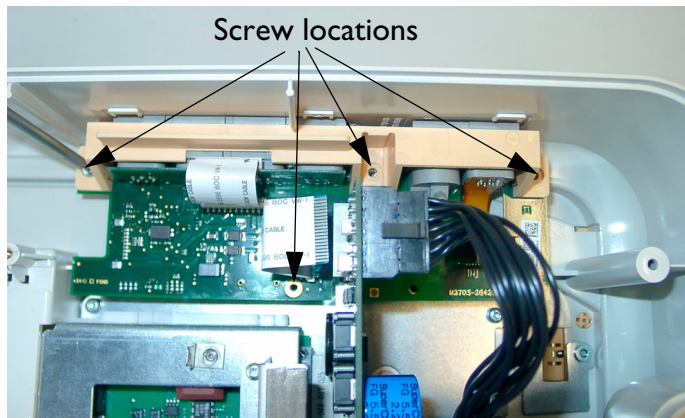
- 1 Remove the top cover assembly (see page 105).
- 2 If the noninvasive blood pressure assembly is fitted, remove it (see page 134).
- 3 If the SpO<sub>2</sub> assembly is fitted, remove it (see page 139). Also remove the SpO<sub>2</sub> board holder by removing the two screws fastening it to the bottom housing.



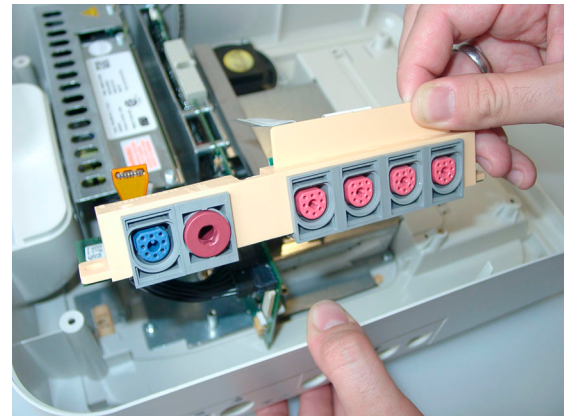
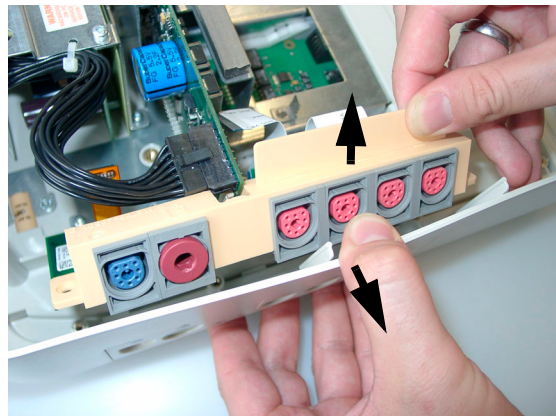
- 4 Disconnect the flat ribbon cable connecting the bus master board to the main CPU board.



- 5 Remove the three screws holding the connector block frame to the bottom housing, and the screw holding the bus master board.



- 6 Lift out the front end assembly. To make the removal easier, apply gentle outward pressure on the bottom housing wall to provide a little more clearance.

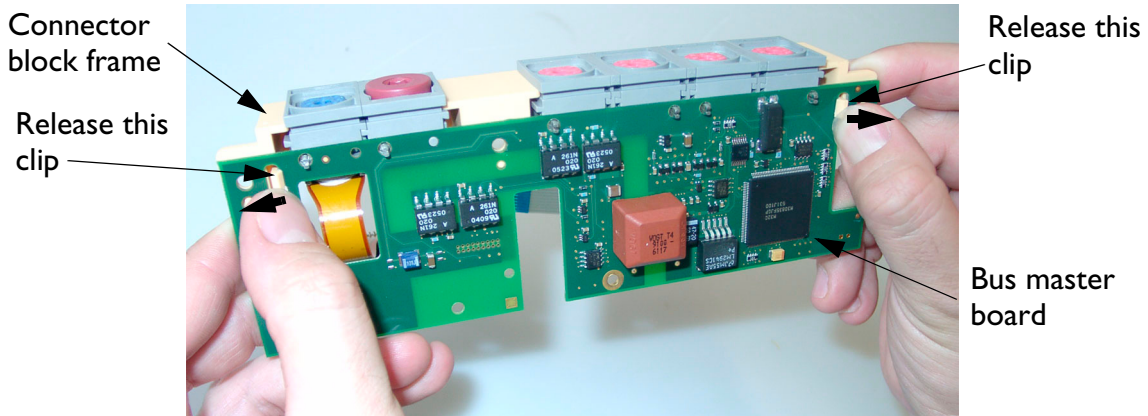


## Replacing the Front End Assembly

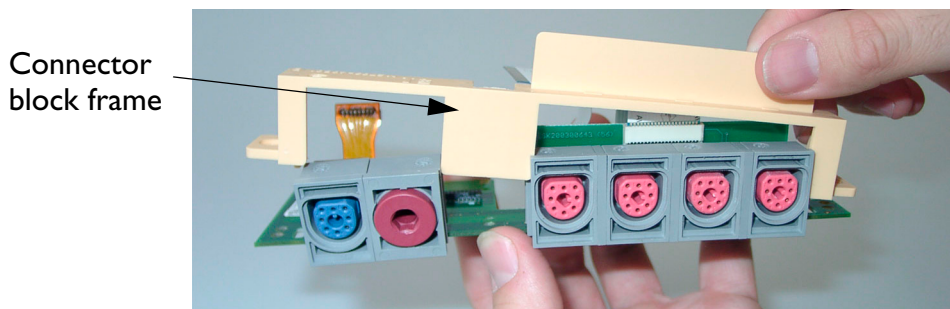
The procedure to replace the front end assembly is a reversal of the removal procedure. Ensure all the cables are properly reconnected.

## Disassembling the Front End Assembly

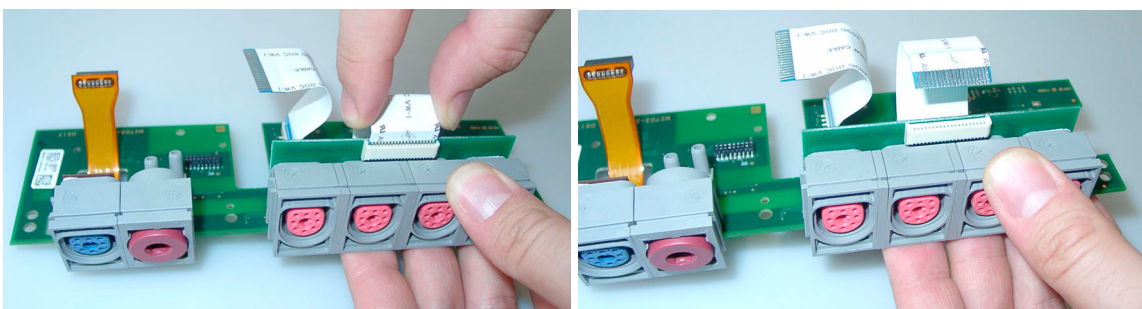
- 1 Remove the front end assembly from the bottom housing (see “Removing the Front End Assembly” on page 143).
- 2 Release the clip at each end of the connector block frame from the slots in the bus master board.



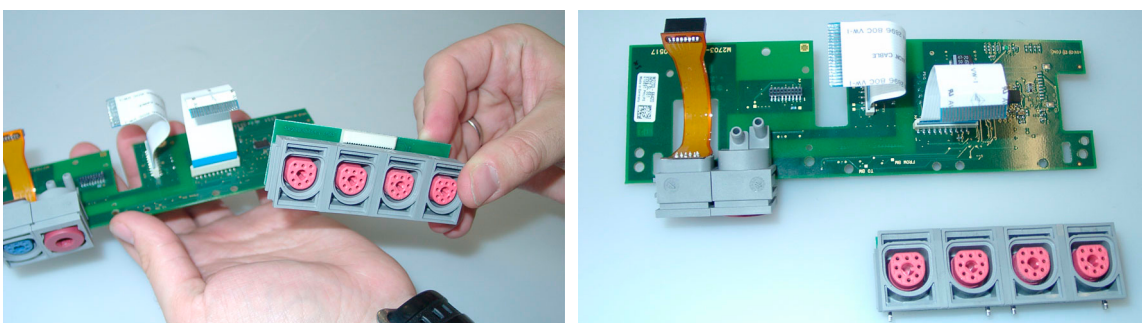
3 Remove the connector block frame.



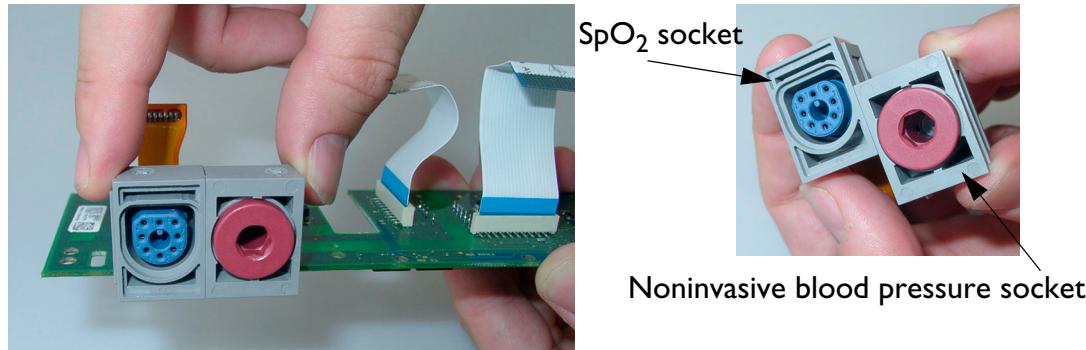
4 Disconnect the flat ribbon cable connecting the fetal sensor socket connector block to the bus master board, then remove the connector block.



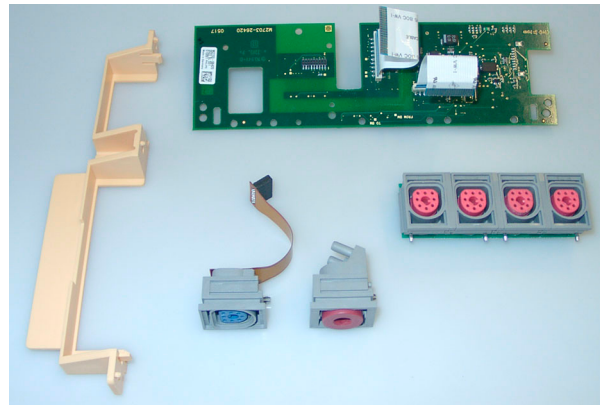
5 Remove the fetal sensor socket connector block.



6 Remove the optional sockets for noninvasive blood pressure and SpO<sub>2</sub>, if fitted. Note that these sockets slide together.



All components of the front end assembly are now separated.



## Reassembling the Front End Assembly

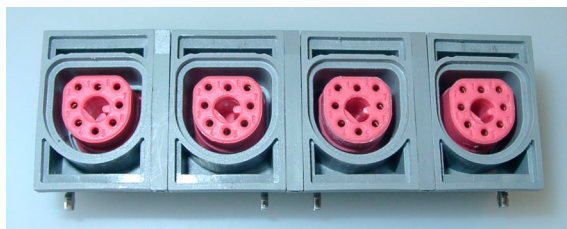
The procedure to reassemble the front end assembly is a reversal of the removal procedure. Ensure all the cables are properly reconnected.

### Exchanging the Bus Master Board

- 1 Remove the front end assembly (see “Removing the Front End Assembly” on page 143).
- 2 Strip the front end assembly down to its component parts (see “Disassembling the Front End Assembly” on page 145).
- 3 Take a new bus master board and reassemble the front end assembly (see “Reassembling the Front End Assembly” on page 147).

### Exchanging the Fetal Socket Connector Block

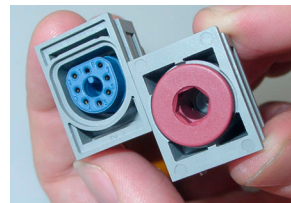
- 1 Remove the front end assembly (see “Removing the Front End Assembly” on page 143).
- 2 Strip the front end assembly down as far as the end of step 5 on page 146 under “Disassembling the Front End Assembly”).
- 3 Take a new fetal socket connector block.



- 4 Reassemble the front end assembly (see “Reassembling the Front End Assembly” on page 147).

## Exchanging the Noninvasive Blood Pressure/SpO<sub>2</sub> Sockets

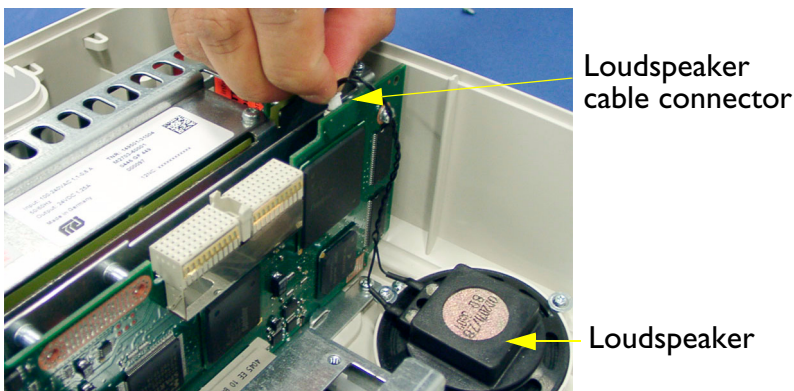
- 1 Remove the front end assembly (see “Removing the Front End Assembly” on page 143).
- 2 Strip the front end assembly down as far as the end of step 6 on page 146 under “Disassembling the Front End Assembly”), but without removing the fetal sensor socket connector block.
- 3 Take a new noninvasive blood pressure and/or SpO<sub>2</sub> socket as applicable. If you need to refit both the noninvasive blood pressure and/or SpO<sub>2</sub> socket, you need to slot them together before attaching them back onto the bus master board.



- 4 Reassemble the front end assembly (see “Reassembling the Front End Assembly” on page 147).

## Exchanging the Loudspeaker

- 1 Remove the top cover assembly (see page 105).
- 2 Disconnect the loudspeaker cable connector from the main CPU board.



- 3 Remove the three screws holding the loudspeaker, and remove the loudspeaker.
- 4 Refit the loudspeaker, making sure to refit the o-ring gasket. If a new gasket is required, it is available as part of the bottom housing small parts kit.

# Transducer Disassembly/Reassembly

This section describes the disassembly and reassembly operations for the transducers.

## Exchanging the Transducer Cable

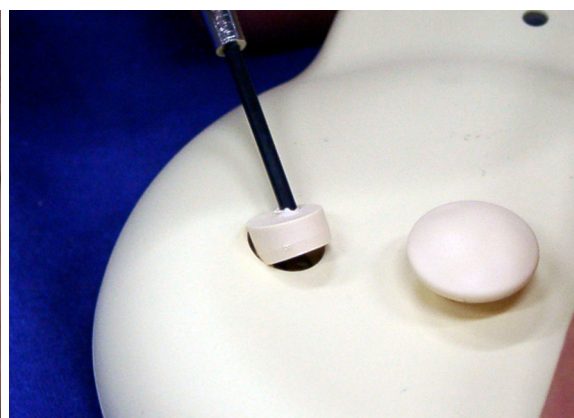
See the “Transducer Cable Assembly” on page 98 for items that come with the cable.

**WARNING** Transducers are calibrated at the factory, and the calibration data for the measurement is stored on the CPU board. Therefore NEVER replace the CPU board with one from another transducer.

**Important when fitting the screw covers!** Do NOT remove the screw covers from the frame to which they are attached. Leave them in place, as it is the only way to align the screw covers correctly. They detach from the frame when you press them into position.

To exchange a transducer cable:

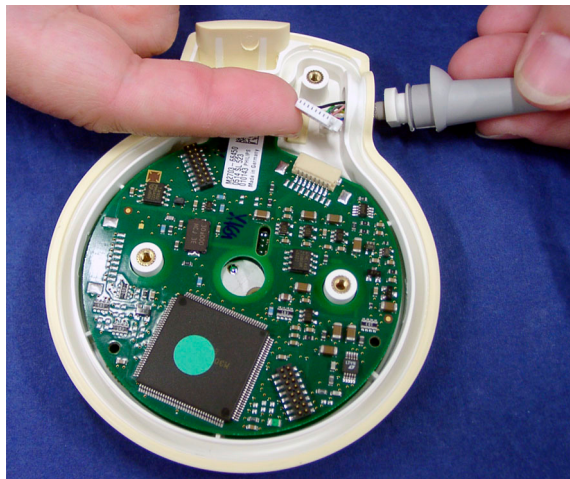
- 1 Pierce a screw cover with a small, flat-bladed screwdriver. **Important!** Do NOT try to prise out a screw cover from the side, without piercing it, as this will damage the transducer top cover.
- 1 Gently rock the screwdriver back and forth until the screw cover comes out. Repeat to remove all three screw covers



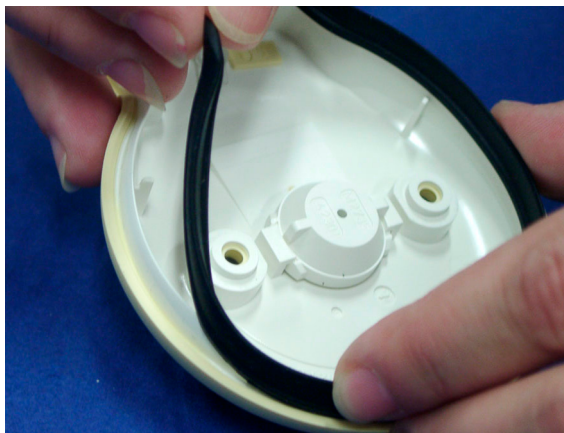
- 2 Remove the three screws, and remove the transducer top cover.



- 3 Disconnect the small cable connector, remove the old cable, and fit the new cable (as a reversal of the removal procedure).



- 4 Remove the sealing gasket from the top cover, and replace it with the new one supplied with the cable. **While handling a Toco/Toco<sup>+</sup> transducer, take care not to displace the strain gauge.** Fit a new gasket to the top cover, ensuring the gasket is properly seated, replace the top cover and secure it with the three screws.



- 5 Leaving the screw covers attached to the frame, carefully align the screw covers with the screw recesses in the top cover. Next, partially press in two of the covers at the same time, then press in the third one (they detach from the frame as you push them in). Then make sure all three covers are pushed completely into the recesses.



## Exchanging the Transducer Belt Button

**CAUTION** NEVER immerse a transducer in liquid if the belt button has been removed, or is loose, broken or damaged.

**M2703-64204**

**Replacement Belt Button Kit**

**Contents:**



**x5**

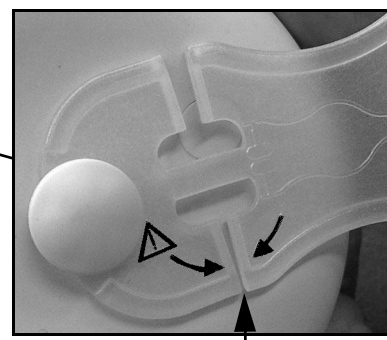
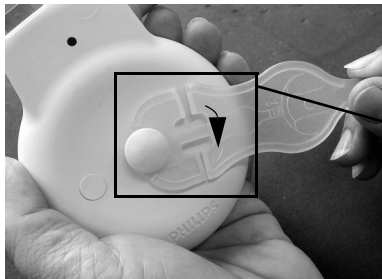


**x1**

- 1 Remove the belt button using the tool provided with the belt button kit.



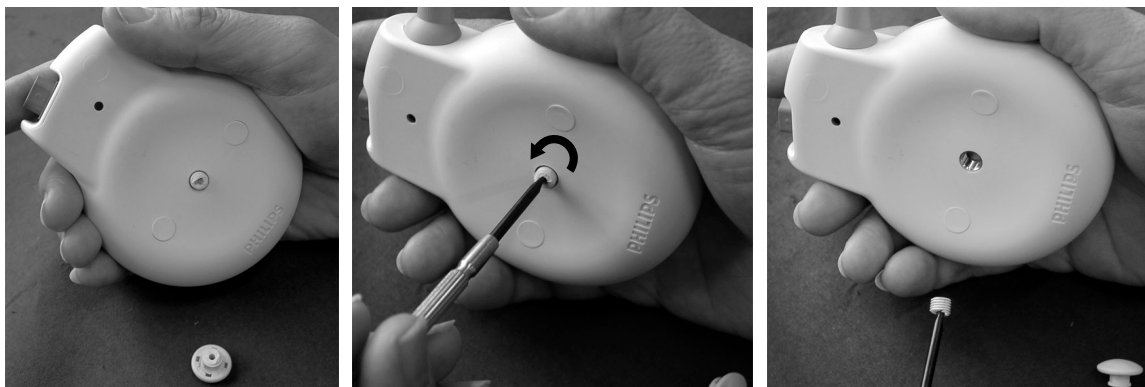
- 2 Dispose of the old belt button. Take a new belt button and fit it to the transducer. Initially, screw the button in by hand about four turns, then complete the job with the supplied tool. Stop applying force when the head of the tool makes contact with the body of the tool at the point indicated by the arrows.



If the belt button is broken:



- 1 Remove the threaded part left in the top cover with a small, flat-bladed screwdriver (2.0-3.0 mm).



2 Then fit a belt button as described on page 151.

# Upgrades

This chapter lists the various upgrade options for the monitors, and describes how to carry out these upgrades.

## FM20/30 Upgrade Options

Upgrade options for the FM20 are prefixed with M2702AU.

Upgrade options for the FM30 are prefixed with M2703AU.

Options B72 and B73 apply to the FM30 only.

FM20/30 Upgrade Options					
Option Number	Option Adds	Parts included with Option			
		Description	Part Number	Contents/Comments	Qty
B71	Noninvasive Blood Pressure (NIBP). No supplies included.	NIBP kit	1253-8416	Connector housing	1
			8120-8912	Ribbon cable	1
			M2703-02301	NIBP assembly holder	1
			M2703-47301	NIBP tubing	1
			M2703-64502	NIBP assembly	1
			M2720-61604	Ground cable	1
			0515-4671	Screw M3x6	5
		<i>Either Connector cover with symbols</i>	M2703-44105	For countries other than USA	1
		<i>Or Connector cover with text</i>	M2703-44115	For USA	1
B72 FM30 ONLY	SpO <sub>2</sub> (only for monitors that already have Noninvasive Blood Pressure installed). No supplies included.	SpO <sub>2</sub> kit	1253-8422	Connector housing	1
			1253-8429	Connector for SpO <sub>2</sub> board	1
			M2703-44108	SpO <sub>2</sub> assembly holder	1
			M2703-66513	SpO <sub>2</sub> board	1
			0515-4671	Screw M3x6	4
		<i>Either Connector cover with symbols</i>	M2703-44106	For countries other than USA	1
		<i>Or Connector cover with text</i>	M2703-44116	For USA	1

FM20/30 Upgrade Options					
Option Number	Option Adds	Parts included with Option			
		Description	Part Number	Contents/Comments	Qty
B73 FM30 ONLY	Noninvasive Blood Pressure and SpO <sub>2</sub> . No supplies included.	NIBP kit	See B71	See B71	See B71
		SpO <sub>2</sub> kit	See B72	See B72	See B72
		<i>Either</i> Connector cover with symbols	M2703-44106	For countries other than USA	1
		<i>Or</i> Connector cover with text	M2703-44116	For USA	1
C73	Triplets monitoring capability.	Customer letter - English	M8000-9399A	Asks customer to contact Philips Support to arrange a software upgrade	1
J22	Dual PS/2 Interface for connecting a keyboard and mouse.	Input device assembly	M8086-67501	“Plug & Play” interface	1
J70	System Interface, 1 × RS232 port and 1 × LAN port.	RS232/LAN I/O card assembly	M2703-67501	“Plug & Play” interface	1
SD0	Rel. D.0 Software Upgrade	Fetal Monitoring Documentation DVD-ROM	M2731-9400C	-	1
		Instructions for Use	M2703-90xxC	Appropriate localized version	1

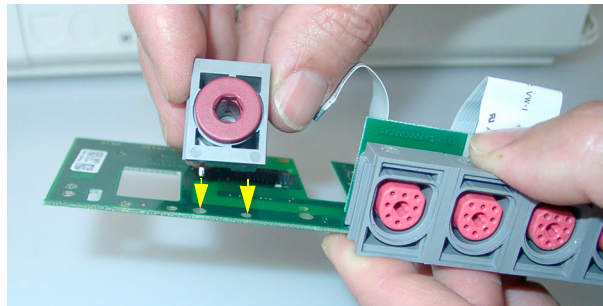
## Installing Upgrade Options

This section covers how to install the options.

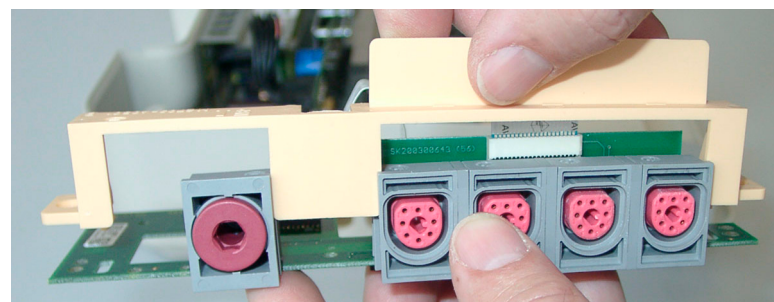
### Option B71

To install the noninvasive blood pressure upgrade:

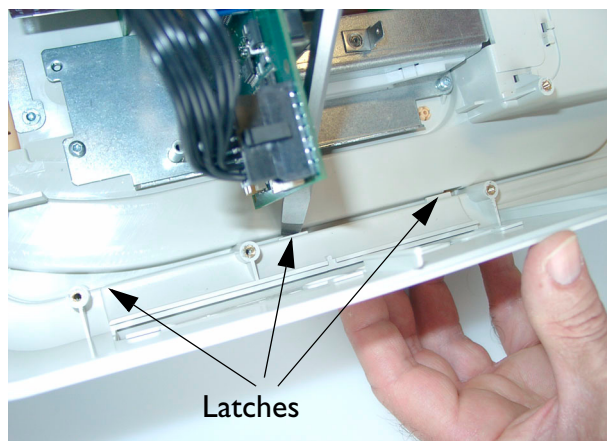
- 1 Remove the top cover assembly as described on page 105.
- 2 Remove the front end assembly (see “Removing the Front End Assembly” on page 143).
- 3 Remove the connector block frame from the front end assembly (see “Disassembling the Front End Assembly” on page 145). Leave the fetal sensor socket connector block in place.
- 4 Take the new noninvasive blood pressure socket. Locate the metal pin and the plastic stud on the underneath of the noninvasive blood pressure socket in the corresponding holes in the bus master board.



5 Replace the connector block frame, making sure that it snaps securely back into the bus master board.



6 To remove the existing connector cover, release the three snap-fit latches holding the connector cover along its bottom edge with a flat-headed screw-driver with a fairly wide blade.



7 Then remove the cover.



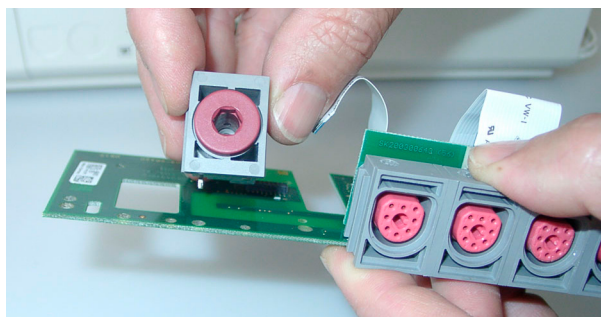
- 8 Fit the new connector cover that comes with the upgrade kit. First, locate the three lugs along the top edge of the connector cover in the corresponding recesses in the bottom housing. Then push the bottom of the cover so that the three latches snap securely into position.



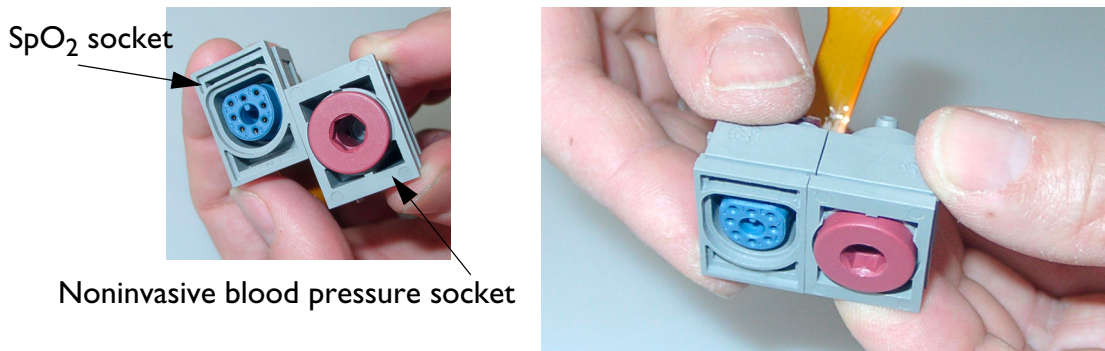
- 9 Replace the front end assembly as described in “Replacing the Front End Assembly” on page 145.
- 10 Fit the noninvasive blood pressure assembly as described in “Replacing the Noninvasive Blood Pressure Assembly” on page 136.
- 11 Replace the top cover assembly as described in “Replacing the Top Cover Assembly” on page 132.
- 12 For tests to perform after upgrading, see “When to Perform Test Blocks” on page 29.

## Option B72 (FM30 only)

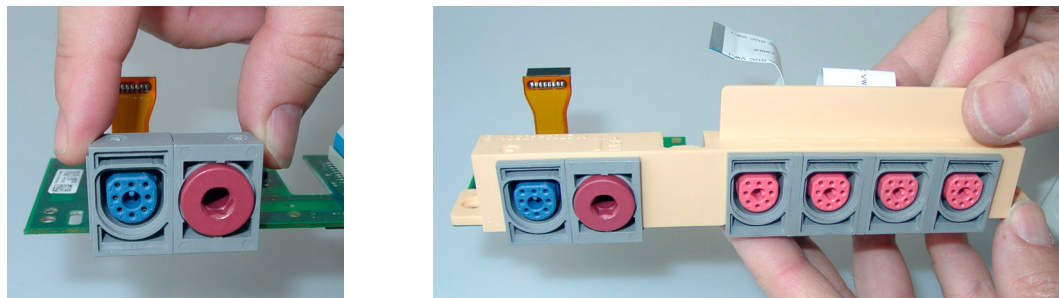
- 1 Remove the top cover assembly as described on page 105.
- 2 Remove the noninvasive blood pressure assembly.
- 3 Remove the front end assembly (see “Removing the Front End Assembly” on page 143).
- 4 Remove the connector block frame from the front end assembly (see “Disassembling the Front End Assembly” on page 145). Leave the fetal sensor socket connector block in place.
- 5 Remove the blood pressure connector from the bus master board.



- 6 Join the new SpO<sub>2</sub> connector to the noninvasive blood pressure connector.



- 7 Locate the metal pins and the plastic studs on the underneath of the noninvasive blood pressure/SpO<sub>2</sub> socket in the corresponding holes in the bus master board. Then replace the connector block frame.



- 8 Remove the existing connector cover as described in step 6 for Option B71 on page 155, then remove the cover.

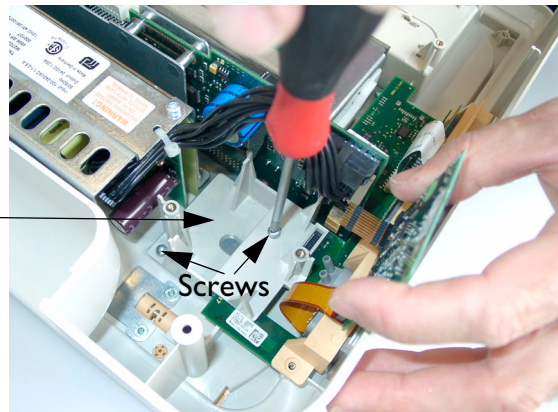


- 9 Fit the new connector cover that comes with the upgrade kit as described in step 8 for Option B71 on page 156.



10 Replace the reassembled front end assembly as described in “Replacing the Front End Assembly” on page 145.

11 Fit the SpO<sub>2</sub> board holder to the bottom housing with two screws from the upgrade kit.



12 Next, fit the SpO<sub>2</sub> board as described in “Replacing the SpO<sub>2</sub> Assembly” on page 141.

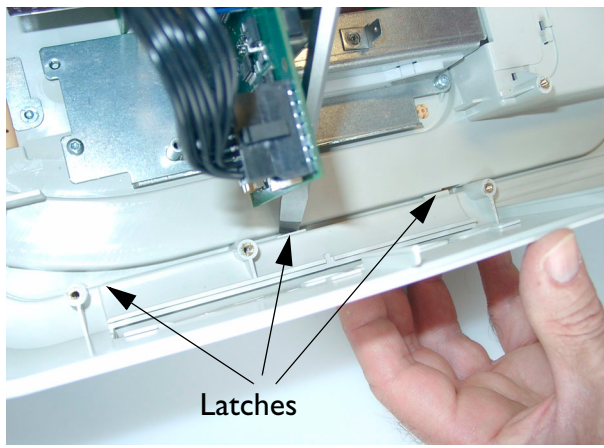
13 Then refit the noninvasive blood pressure assembly as described in “Replacing the Noninvasive Blood Pressure Assembly” on page 136.

14 Replace the top cover assembly as described in “Replacing the Top Cover Assembly” on page 132.

15 For tests to perform after upgrading, see “When to Perform Test Blocks” on page 29.

## Option B73 (FM30 only)

- 1 Follow steps 1 to 3 inclusive for Option B71.
- 2 Then follow steps 6 and 7 for Option B72.
- 3 To remove the existing connector cover, release the three snap-fit latches holding the connector cover at the bottom with a flat-headed screw-driver with a fairly wide blade.



- 4 Fit the new connector cover that comes with the upgrade kit as described in step 8 for Option B71 on page 156.



- 5 Follow steps 9 to 14 inclusive for Option B72.
- 6 For tests to perform after upgrading, see “When to Perform Test Blocks” on page 29.

## Option C73

You enable this feature upgrade to triplets capability using the Support Tool. Refer to the *Support Tool Instructions for Use* for details of the upgrade procedure.

## Options J22 and J70

The interfaces require no special upgrade procedures as they are “plug and play” boards.

## Option SD0

You can upgrade the monitor software to Rel. D.0 using the Support Tool. See “Software and Firmware Upgrades” on page 159.

# Software and Firmware Upgrades

The software of the monitor and the firmware of the transducers and other system components can be upgraded by a software download from a PC running the Support Tool. You connect the monitor to the PC via a LAN connection. You need:

- Industry standard PC
- Support Tool
- LAN / RS232 system interface
- LAN interface cable for the Support Tool

Several Avalon fetal monitors can be upgraded in parallel with the Support Tool. All monitors in an installation can be upgraded at once, if desired.

The transducers can be upgraded one at a time, even though more than one may be plugged into the monitor at the same time.

**When upgrading to a new monitor software revision, we recommend that you check that all system hardware components have the latest firmware revision, and upgrade these if necessary.**

- NOTE** Make sure that there is no interruption of the LAN or the power connection during the upgrade procedure. Refer to the *Support Tool Instructions for Use* for details of the upgrade procedure. Contact Philips Support for further details.

For tests to perform after upgrading, see “When to Perform Test Blocks” on page 29.

# Understanding Configuration

This chapter, together with the Settings appendix, is for anyone making permanent changes to the configuration of an Avalon Fetal Monitor. You must understand English, be familiar with the monitor and its *Instructions for Use*, know how to make changes to measurements and settings in Monitoring Mode, and understand the clinical implications of any changes you make.

**WARNING** Changing the configuration may alter the way the monitor performs when monitoring patients. Do not change anything unless you are aware of the possible consequences, especially if you are monitoring a patient while in Configuration Mode.

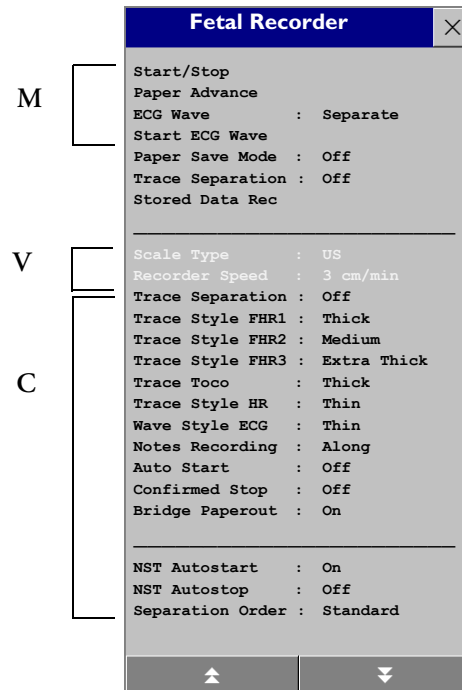
## What is Configuration Mode?

Configuration Mode is a password-protected operating mode that lets expert users make permanent changes to the monitor configuration. It is an extension of Monitoring Mode; it contains all of the settings available in Monitoring Mode plus the settings that are accessible only in Configuration Mode.

For example, the **Fetal Recorder** menu accessible in Monitoring Mode contains the settings marked M (not necessarily in exact order).

Items that are visible in Monitoring Mode but that can only be changed in Configuration Mode are marked V.

In Configuration Mode you can change the additional settings marked C.



Configuration Mode is a password-protected operating mode that lets expert users make permanent changes to the monitor configuration. It is an extension of Monitoring Mode; it contains all of the settings available in Monitoring Mode plus the settings that are accessible only in Configuration Mode.

In Monitoring Mode, you do not see all possible menu entries. There are two types of menu entry you will see:

- Items in black text are those items you can operate or access to change the setting.
- Items that are grayed out are those items that you can see but not operate or access to change the setting.

## Understanding Settings

You can change two main categories of settings in Configuration Mode: Global Settings, and the monitor and measurement settings stored in User Defaults. The monitor ships with preset configurations for Global Settings and the Factory Default settings that are suitable for common monitoring situations. This guide tells you how to develop your own configurations.

**Active Settings** are the current settings the monitor uses, including any adjustments made by the last user. Active Settings are not permanent, but are retained after a loss of mains power.

**The User Defaults** is a complete configuration of monitor and measurement settings blocks stored in the monitor's long-term memory. You can change individual settings and store them in the User Defaults. In other words, you can store the Active Settings, modified to your preference, in the User Defaults. Alternatively, you can load a complete configuration (taken from another monitor, for example) into the User Default from the Support Tool. The User Defaults is the user's preferred configuration, and these personalized settings can be restored by loading the User Defaults.

Following a patient discharge, or if the monitor was turned off for more than one minute, the User Defaults is automatically loaded if **Automat. Default** is set to **Yes**.

**The Factory Default** is a complete configuration predefined at the factory. You cannot modify it. In Configuration mode, you can load the Factory Default as the Active Settings. You can use the Factory Default as the basis for producing your own User Defaults.

**Global Settings** are typically set once at monitor installation by service personnel and include settings such as **Line Frequency**, or **QRS Type**. Global settings are independent of the User Defaults, so when you load the Factory Defaults, Global Settings remain as they were. They can only be changed in Configuration Mode and are automatically stored to the monitor's permanent memory with each change. Global settings can be cloned.

**Hardware Settings** are typically set once at monitor installation by service personnel. Most hardware settings can only be changed in Service mode. Hardware settings include settings such as **Keyboard** layout or the **Intensity** setting for the thermal printhead. Like Global settings, they are independent of the User Defaults, and any changes you make to the Hardware Settings configuration are automatically stored, there is no need to save them in an extra step. Unlike Global Settings, hardware settings must be entered for each monitor individually, because they **cannot be cloned**.

## Entering and Leaving Configuration Mode

Only people authorized to do so by their institution should make changes in Configuration Mode. They require the Configuration password.

Switching between Monitoring and Configuration Mode does not affect the active settings. You can even continue to monitor patients while in Configuration Mode. The password for Configuration Mode is given in Chapter 1.

To enter Configuration Mode:

- 1 In the **Main Setup** menu, select **Operating Modes**.
- 2 Select **Config** and enter the password.

The monitor displays **Config** in the center of the Screen while you are in Configuration Mode.

Before you leave Configuration Mode, always be sure to store any changes you made. You must store the changes you made in the User Default.

To leave Configuration Mode either:

- ◆ In the **Main Setup** menu, select **Operating Modes** and then select the operating mode you require or
- ◆ Switch the monitor off, then switch it on again.
  - If you switch the monitor off and then on again after less than one minute, it returns in Monitoring Mode with the same settings (“hotstart”).
  - If you leave the monitor switched off for more than one minute, the User Default is loaded when you switch back on if **Automat. Default** is set to **Yes**.

## Storing Changes in the User Defaults

You can load a complete configuration for the monitor via the Support Tool, or you can change individual settings within the Active Settings. The monitor remembers any changes made when you switch between Monitoring Mode and Configuration Mode. The changes made in Configuration Mode can be stored permanently in the User Defaults.

- 1 Make the changes to the individual measurements or monitor settings.

- 2 Select the **Defaults** SmartKey  .

- 3 Select **Store Defaults** from the pop-up keys at the bottom of the screen.

Load Defaults	Store Defaults		Factory Default
------------------	-------------------	--	--------------------

- 4 Select Confirm to store the settings in the User Default.

<b>To store the current settings as user defaults select Confirm</b>	<b>Confirm</b>	<b>Cancel</b>
--	----------------	---------------

Be aware that if you don't store changes they will be reset to the monitor's stored configuration when you:

- change from Configuration or Monitoring Mode to Service or Demonstration Mode.
- switch off the monitor for more than one minute and **Automat. Default** is set to **Yes**.

## Loading the Factory Default

Load the Factory Default to restore the Active Settings to those that were set at the factory and shipped with the monitor. You may want to do this to fall back to a known, reliable configuration, or you may want to use the default settings as a basis for making your own customized settings that you would save as the User Defaults.

To load the Factory Defaults:

1 Select the **Defaults** SmartKey  .

2 Select **Factory Default** from the pop-up keys at the bottom of the screen.

<b>Load Defaults</b>	<b>Store Defaults</b>		<b>Factory Default</b>
--------------------------	---------------------------	--	----------------------------

3 Select Confirm to load the settings stored in the User Defaults.

<b>To reset the current settings to the factory defaults select Confirm</b>	<b>Confirm</b>	<b>Cancel</b>
---	----------------	---------------

4 Check the paper scale type setting after loading the Factory Default, and change it if necessary.

When you load the Factory Default, note that:

- Global Settings are not reset.
- Hardware Settings are not reset.

## Loading the User Defaults

Load the User Defaults to restore the Active Settings to the stored customized settings:

1 Select the **Defaults** SmartKey  .

2 Select **Load Defaults** from the pop-up keys at the bottom of the screen.

<b>Load Defaults</b>	<b>Store Defaults</b>		<b>Factory Default</b>
--------------------------	---------------------------	--	----------------------------

3 Select Confirm to load the settings stored in the User Defaults.

<b>To reload the user default settings select Confirm</b>	<b>Confirm</b>	<b>Cancel</b>
---	----------------	---------------

When you load the User Defaults, note that:

- Global Settings are not reset.
- Hardware Settings are not reset.

# Loading Configurations Using the Support Tool

Use of the Support Tool is restricted to technical personnel who have been trained in its use by Philips.

Using the Support Tool you can clone multiple monitor configurations and store the configuration file in a format that can be e-mailed.

The Support Tool lets you make a backup of your configuration and any changes you make. See the *Support Tool Instructions for Use* for details about storing, cloning, and maintaining your configurations.

If you make a lot of configuration changes to monitors throughout your institution, you are strongly advised to acquire the Support Tool so that you can backup this work and restore configurations if necessary.

## About Configuration Files (.cfg)

Each.cfg file contains all the settings saved in a configuration. These are complete configurations including all measurement and monitor settings. Files of the format.cfg can only be read and modified using the Support Tool. A checksum protects the contents of the configuration files, checking for example whether files were corrupted during e-mail transfer. Corrupted files will be rejected by the Support Tool.

There are two kinds of configurations:

- **initial configurations** are configurations provided by the factory. Each initial configuration supports all languages that the monitor is currently shipped with. Initial configuration files cannot be modified using the Support Tool. When an initial configuration is cloned to a monitor, the configuration is automatically adjusted to incorporate some monitor-specific attributes, for example, the language and product options. Cloning this configuration from the monitor back to the Support Tool changes it to a single-language user configuration that can then be modified using the Support Tool.
- **user configurations** are configurations that can be edited, deleted, or added to using the Support Tool. They can either be copied from a monitor or from a configuration stored with the Support Tool files on your computer. As user configurations are language dependent, always use a configuration taken from a monitor with the correct language. If you clone a user configuration to a monitor with a different language, all user adjustable texts are reset to factory defaults the first time you switch the monitor on.

## Selecting the Correct Configuration

When cloning configurations, always use a configuration designed for the target device, and with the same options for application area (Hxx Option) and number of waves (Axx Option).

This is an example of an Avalon configuration file:

**H70 A01, 60Hz, Scale 240, VGA, FM30-30, initial, F.0X.xx, Rev  
xxx.cfg**

The name of a configuration file consists of codes to identify, where appropriate:

- the **Hxx** (application area) option and **Axx** (wave number) it is optimized for.
- the line frequency (50Hz or 60Hz).

- the paper scale type.
- the resolution of the majority of screens supplied with the config file.
- the monitor model (**FM30** **and** **FM30** in our example) that the config file is optimized for.
- the word “**initial**” to mark an initial configuration provided by the factory.
- the software revision of the product it is optimized for. The letter “x” is a placeholder for any number from 0 to 9.
- the revision code used to track changes during the configuration creation process (only the latest revision is bundled with the tool).

# Configuration Settings Appendix

The monitor is pre-configured with factory defaults settings when it is shipped. This section documents these factory default configuration settings. If you change the User Defaults, this document will no longer reflect your configuration, so you must note any changes you make in the editable version of this appendix provided on the documentation DVD-ROM. The initial configuration of your monitor may vary slightly depending on your geography and on the options purchased.

In most cases, there is one set of factory default settings listed in the tables under “Factory Defaults”. Where there is more than one set of defaults (due to geography-specific options, for example), these are noted in the tables. The tables contain a blank section called “User Defaults”, where you can document your preferred, customized settings saved to the User Defaults.

## Documenting Monitor Configurations

To help you document your monitor’s configuration, the configuration tables from this appendix are also provided as a Word document on the documentation DVD-ROM supplied with the monitor. To document the configurations you create, edit this document using a word-processing program to reflect the configuration and then save it under an appropriate name.

As Philips cannot take responsibility for changes made to this document in the \*.doc format, you must only use the.pdf version of this appendix as a reference for the initial configuration settings supplied with the monitor.

The configuration implications are only provided in the.pdf version of this appendix. You must read this document before you modify monitor configurations.

\*Word is a registered trademark of the Microsoft Corporation.

## Using the Configuration Tables

The “breadcrumb trail” at the top of each table indicates the path you should follow to access the settings in the table: in this example, to configure the fetal recorder settings, in the **Main Setup** menu, select **Measurements** and then select **Fetal Recorder**.

## Configuration Table Example

This is a (shortened) example of a configuration table, as you will find it in the following sections of this manual.

Item Name	Factory Defaults	Choice	User Defaults
Scale Type	Geography-specific	US (scale = 30-240) Internat'l (scale = 50-210)	
Recorder Speed	3 cm/min	1, 2, 3 cm/min	
Trace Style FHR1	Thick		
Trace Style FHR2	Medium		
Trace Style FHR3	Extra Thick	Thin, Medium, Thick, Extra Thick	
Trace Style HR	Thin		
Wave Style ECG	Thin		
ECG Wave	Separate	Separate, Overlap	
Auto Start	Off	Off, On	

**Item Name** The leftmost column in each table lists the individual configuration items. These items correspond to the menu items in the relevant setup menu in the monitor.

**Factory Defaults** This section deals with the factory default settings for each configuration item.

**Choice** This lists the possible choices for the settings you can configure.

**User Defaults** In each table, columns are left blank for you to enter the settings you change.

**NOTE** You cannot print out the configuration from the monitor: these tables are your only documentation of the configuration you implement for each monitor. We strongly recommend that you always write down any changes you make and keep this record safely.

## Understanding Configuration Implications

When you permanently change any element of the configuration, you must consider the effect of the new configuration on both patient and application behavior. For more information on the context of the configuration settings, see the monitor *Instructions for Use*. Always ensure that the monitor users are aware of the configuration settings.

# Measurement-Related Settings

This section lists all the measurement-related settings. They define how the monitor measures patient data. Document the settings you configure in the empty columns.

Read any information on Configuration Implications at the end of the sections before you make any configuration changes.

## Color Configuration

The color setting for each measurement defines the color for its numeric (and wave, if applicable). The color setting for Pulse is taken from the active pulse source. The choice for color is: **Red, Green, Yellow, Blue, Magenta, Cyan, White, Pink, Orange, Light Green, Light Red**

## Configuring FHR (Ultrasound)

**Main Setup --> Measurements --> FHR (1/2/3)**

Item Name	Factory Defaults	Choice	User Defaults
FHR Sound Volume	6	0..10	
High Limit	150 bpm	70..210 bpm, in 10 bpm steps	
Low Limit	110 bpm	60..200 bpm, in 10 bpm steps	
Alarms	On	Off, On	
Fetal Movement	On	Off, On	
High Delay	60 sec	10..300 sec, in 10 second steps	
Low Delay	60 sec	10..300 sec, in 10 second steps	
SignalLoss Delay	60 sec	10..300 sec, in 10 second steps	
Color	Orange	See "Color Configuration" on page 169	

## FHR Configuration Implications

**High Limit/Low Limit, High Delay/Low Delay, SignalLoss Delay** All FHRs, including DECG, share the same alarm limits and delays, and can be set from any FHR channel.

**Alarms** This lets you switch **On** FHR alarms. Your monitor must be configured to alarm mode **All** to enable the FHR alarms.

**Fetal Movement** Fetal movement profile can be enabled from any FHR channel, even though the fetal movement detection itself only applies to FHR1.

## Configuring Toco

Main Setup --> Measurements --> Toco

Item Name	Factory Defaults	Choice	User Defaults
Toco Gain	100%	50%, 100%	
Color	Green	See “Color Configuration” on page 169	

## Configuring IUP

Main Setup --> Measurements --> IUP

Item Name	Factory Defaults	Choice	User Defaults
IUP Unit	mmHg	mmHg, kPa	
Color	Green	See “Color Configuration” on page 169	

## Configuring DFHR (DECG)

Main Setup --> Measurements --> DFHR

Item Name	Factory Defaults	Choice	User Defaults
FHR Sound Volume	6	0..10	
High Limit	150 bpm	70..210 bpm, in 10 bpm steps	
Low Limit	110 bpm	60..200 bpm, in 10 bpm steps	
Alarms	On	Off, On	
ArtifactSuppress	On	Off, On	
High Delay	60 sec	10..300 sec, in 10 second steps	
Low Delay	60 sec	10..300 sec, in 10 second steps	
SignalLoss Delay	60 sec	10..300 sec, in 10 second steps	
Color	Orange	See “Color Configuration” on page 169	
ECG Wave	Off	On/Off	

### DFHR Configuration Implications

Your monitor must be configured to alarm mode **All** to enable the FHR alarms.

**High Limit/Low Limit, High Delay/Low Delay, SignalLoss Delay** All FHRs, including DECG, share the same alarm limits and delays, and can be set from any FHR channel.

**Alarms** This lets you switch **On** FHR alarms.

**ArtifactSuppress** This lets you switch artifact suppression **On** (artifacts are suppressed) and **Off** (no artifact suppression: use this setting if you suspect fetal arrhythmia).

## Configuring MHR (ECG)/Pulse

Main Setup --> Measurements --> ECG

Main Setup --> Measurements --> Pulse

Item Name	Factory Defaults	Choice	User Defaults
High Limit	120 bpm	31..300 bpm in steps of 1 bpm (31 to 40 bpm) in steps of 5 bpm (40 to 300 bpm)	
Low Limit	50 bpm	30..295 bpm in steps of 1 bpm (30 to 40 bpm) in steps of 5 bpm (40 to 295)	
Alarms	On	Off, On	
QRS Volume	1	0..10	
Δ ExtrTachy	20 bpm	0..50 bpm, in steps of 5 bpm	
Tachy Clamp	200 bpm	150..240 bpm, in steps of 5 bpm	
Δ ExtrBrady	20 bpm	0..50 bpm, in steps of 5 bpm	
Brady Clamp	40 bpm	30..100 bpm, in steps of 5 bpm	
Color	Red (ECG) Cyan (Pulse from SpO <sub>2</sub> )	See "Color Configuration" on page 169	

## ECG/Pulse Configuration Implications

Your monitor must be configured to alarm mode **All** to enable the MHR alarms.

**High Limit/Low Limit** MHR (MECG) and Pulse share the same alarm limits. These alarm limits apply to the current alarm source, either HR or Pulse. Note that if you change the High/Low alarm limits in the Setup ECG menu, this will also change the High/Low alarm limits in the Setup Pulse menu and vice versa.

**Alarms** This lets you switch **Off** HR alarms. If you change the **Alarms** setting in the Setup ECG menu, this will also change the **Alarms** setting in the Setup Pulse menu and vice versa.

**$\Delta$  ExtrTachy,  $\Delta$  ExtrBrady** Extreme bradycardia and extreme tachycardia alarms are based on the HR/Pulse limit alarms. In Configuration Mode, you use the  $\Delta$  **ExtrTachy** and  $\Delta$  **ExtrBrady** setting to define the difference between the heart rate limit and the extreme limit. For example, if the heart rate high limit is 120 bpm and the difference is 20 bpm then the extreme tachycardia limit is 140. HR and Pulse share the same alarm limits. The  $\Delta$  **ExtrTachy** and  $\Delta$  **ExtrBrady** settings apply to the current alarm source, either HR or Pulse. If you change the  $\Delta$  **ExtrTachy** or  $\Delta$  **ExtrBrady** setting in the Setup ECG menu, this will also change the  $\Delta$  **ExtrTachy** or  $\Delta$  **ExtrBrady** setting in the Setup Pulse menu and vice versa.

**Tachy Clamp, Brady Clamp** The Brady and Tachy clamp allows you to configure a safety threshold for the extreme bradycardia and tachycardia alarm limits. For example, if the low heart rate limit is 50 bpm and the  $\Delta$  **ExtrBrady** setting is 20 bpm ( $50 \text{ bpm} - 20 \text{ bpm} = 30$ ) with a Brady clamp set at 40, the resulting extreme bradycardia limit would be 40 bpm (instead of 30 bpm). If the clinician sets the HR alarm limit above or below the limit clamps for an individual patient, the limit clamps become the extreme brady or extreme tachy alarm (these are red alarms). Be sure to set the clamps beyond the configured HR limits.

HR and Pulse share the same alarm limits. The **Tachy Clamp** and **Brady Clamp** settings apply to the current alarm source, either HR or Pulse. If you change the **Tachy Clamp** or **Brady Clamp** setting in the Setup ECG menu, this will also change the **Tachy Clamp** or **Brady Clamp** setting in the Setup Pulse menu and vice versa.

**Alarms Off** Note that changing the **Alarms Off** setting in the Setup ECG menu also changes the **Alarms Off** setting in the Setup Pulse menu and vice versa.

## Configuring SpO<sub>2</sub>

Main Setup --> Measurements --> SpO<sub>2</sub>

Item Name	Factory Defaults	Choice	User Defaults
High Limit	100	51..100 bpm, in 1 bpm steps	
Low Limit	90	50..99 bpm, in 1 bpm steps	
Desat Limit	80	50..99 bpm, in 1 bpm steps	
Alarms	On	Off, On	
QRS Volume	1	0..10	
Tone Modulation	Yes	Yes, No	
Tone Mod. Type	Enhanced	Enhanced, Standard	
Average	10 sec	20, 10, 5 sec	
High Alarm Delay	10 sec	0..30 sec, in 1 second steps	
Low Alarm Delay	10 sec	0..30 sec, in 1 second steps	
Desat Alarm Delay	20 sec	0..30 sec, in 1 second steps	
NBP Alarm Suppr.	On	Off, On	
Color	Cyan	See "Color Configuration" on page 169	

### SpO<sub>2</sub> Configuration Implications

**SpO<sub>2</sub>** The **On/Off** state of the SpO<sub>2</sub> measurement cannot be preconfigured. SpO<sub>2</sub> is automatically switched **On** when an SpO<sub>2</sub> sensor is connected to the monitor.

**Average** The SpO<sub>2</sub> numeric represents an average value calculated from the sum of SpO<sub>2</sub> values measured during the averaging time. **Average** lets you adjust the averaging time between **5**, **10**, and **20** seconds.

**High/Low/Desat Alarm Delay** The alarm delay defines the amount of time that the averaged SpO<sub>2</sub> value needs to be above or below the corresponding alarm limits before an alarm is activated.

**NBP Alarm Suppr.** Set **NBP Alarm Suppr.** to **On** to suppress INOPs that would otherwise be generated when you measure NBP on the same limb as SpO<sub>2</sub>. If **NBP Alarm Suppr.** is configured to **On**, the monitor automatically remembers the SpO<sub>2</sub> value measured before cuff inflation and suppresses any SpO<sub>2</sub> INOPs while the cuff is inflated.

## Configuring Noninvasive Blood Pressure (NBP)

Main Setup --> Measurements --> NBP

Item Name	Factory Defaults	Choice	User Defaults
Pulse(NBP)	On	Off, On	
Alarms from	Systolic	Sys., Dia., Mean, Sys & Dia, Dia & Mean, Sys & Mean, Sys&Dia&Mean	
Sys. High	160	95..270 mmHg	
Sys. Low	90	30..155 mmHg	
Dia. High	90	55..245 mmHg	
Dia. Low	50	10..85 mmHg	
Mean High	110	65..255 mmHg	
Mean low	60	20..105 mmHg	
Alarms	On	Off, On	
NBP	On	Off, On	
Repetition Time	15 min	1, 2, 2.5, 3, 5, 10, 15, 20, 30, 45, 60, 120 min	
Auto/Manual	Manual	Auto/Manual	
Unit	mmHg	mmHg, kPa	
Done Tone	Off	Off, On	
Start Time	Synchronized	Synchronized, NotSynchron.	
VP Pressure	60 mmHg	20..120 mmHg in 5 mmHg steps	
Reference	Auscultatory	Auscultatory, Invasive	
NBP Time	Meas Time	Meas Time, Next Meas	
Color	Red	See "Color Configuration" on page 169	
Closevalves <sup>1</sup>	Off	Off, On	

1.Service Mode only.

### NBP Configuration Implications

**Start Time** If you set **Start Time** to **Synchronized**, the monitor will time the second measurement in a series to coincide with the next easy-to-document time. For example, if you start the first measurement at 08:23, and the **Repetition Time** is set to 10 minutes, the monitor will automatically perform the next measurement at 8:30, then 8:40 and so on.

**Done Tone** Set **Done Tone** to **On** if you want to hear a short prompt tone at completion of each NBP measurement.

**VP Pressure** This setting determines the cuff pressure used during a Veni Puncture inflation. The cuff deflates automatically after a set time (170 seconds) if it is not manually deflated beforehand.

**Reference** The NBP measurement reference method can be **Auscultatory** or **Invasive**. **Invasive** delivers NBP values that very closely approximate values measured intra-arterially. **Auscultatory** delivers NBP values that very closely approximate values measured using the manual cuff method. The two references can exhibit a difference of 20 to 30 mmHg in patients with elevated pressures, with the auscultatory reference registering the lower values.

## Monitor-Related Settings

This section lists all the monitor-related settings (anything other than measurements). Read any information on Configuration Implications at the end of the relevant tables before you make any configuration changes.

### Configuring Alarms

Main Setup --> Alarms --> Alarm Settings

Item Name	Factory Defaults	Choice	User Defaults
Alarm Volume	5	0..10	
Alarms Off	3 min	1, 2, 3 min, infinite	
Visual Latching	Red&Yell	Red&Yell, Red Only, Off	
Audible Latching	Red&Yell	Red&Yell, Red Only, Off	
Alarm Sounds	Traditional	Traditional, ISO	
Alarm Low	4	0..10	
Alarm Text	Standard	Standard, Extended	
Alarm Mode	INOP Only	All, INOP Only	

### Alarm Settings Configuration Implications

**Alarm Volume** Use this setting to define the base volume of the red and yellow audible alarm indicators and the INOP tones.

**Alarms Off** Use this setting to determine how long the monitor's alarm capabilities will be switched off when the user selects the **Alarms Off** or **Pause Alarms** key. Possible choices are: **1min**, **2min**, **3min**, **Infinite**. Be aware that if you configure **Alarms Off** to **Infinite**, all of the monitor's alarming capabilities will be permanently switched off when the user selects the **Alarms Off** key.

**Alarm Low** Use this setting to define a minimum value for the alarm volume. The alarm volume cannot be set lower than this value.

**Alarm Sounds** Use this setting to change the alarm sound of the monitor to suit the alarm standards valid in your hospital.

- **Traditional**: The traditional (“Carenet”) sounds used in previous HP/Agilent/Philips patient monitor generations.
- **ISO**: A new set of alarm sounds that complies with the ISO/IEC Standard 9703-2.

**Alarm Text** Use this setting to define how alarm messages are presented on the monitor screen:

- **Standard**: Alarm texts are displayed in text form, for example **\*\* FHR1 LOW**

- **Extended:** Alarm texts are displayed as numeric values, for example, \*\* **FHR1 94 < 110**, where the second number shows the current alarm limit, and the first number shows the maximum amount by which this limit was exceeded.

**Alarm Mode** There are possible alarm modes for the monitor:

- **All:** Patient alarms and INOPs are enabled, with all audible and visual indicators active.
- **INOP only:** Only INOPs are enabled, with audible and visual indication active. This is the default alarm mode. Note that in **INOP only** mode, no patient alarms are enabled or indicated. No alarm limits or alarm off icons are displayed. No patient alarm settings are available in the setup menus.

## Configuring the NST Timer

Main Setup --> NST Timer

Item Name	Factory Default	Choice	User Default
Run Time	20 min	10..60 minutes, in increments of 5 minutes	
Notification	Sound	Alarm, Sound, No Sound	
Timer	Off	Off, On	
Timer Volume	4	0..10	

### NST Timer Configuration Implications

**Run Time** The run time can be set between 10 and 60 minutes.

**Notification** When the NST timer expires, its color changes from blue to green, and a message appears in the monitor status line on the Main Screen. The setting **Notification** lets you configure an alarm or a single tone as additional means of notification:

- Select **Alarm** to receive an INOP alarm when the timer expires.
- Select **Sound** to hear a single tone when the timer expires.
- Select **No Sound** for no additional notification.

**Timer** The timer is not displayed on the screen as default.

## Configuring Fetal Recorder Settings

Main Setup --> Fetal Recorder

Item Name	Factory Defaults	Choice	User Defaults
Scale Type	Geography-specific	US (scale = 30-240) Internat'l (scale = 50-210)	
Recorder Speed	3 cm/min	1, 2, 3 cm/min	
Trace Style FHR1	Thick		
Trace Style FHR2	Medium		
Trace Style FHR3	Extra Thick	Thin, Medium, Thick, Extra Thick	
Trace Style Toco	Thick		
Trace Style HR	Thin		
Wave Style ECG	Thin		
ECG Wave	Separate	Separate, Overlap	
Notes Recording	Along	Along, Across	
Auto Start	Off	Off, On	
Confirmed Stop	Off	Off, On	
Bridge Paperout	On	Off, On	
NST Autostart	On	Off, On	
NST Autostop	Off	Off, On	
Paper Save Mode	Off	Off, On	
Trace Separation	Off	Off, On	
Separation Order	Standard	Standard, Classic	
Cal. Offset <sup>1</sup>	Calibrated at factory	0=right, 10=left	
Intensity <sup>1</sup>	Set at factory	1..5	

1. Can be changed in Service Mode only

## Recorder Configuration Implications

**Scale Type** The initial setting depends on the geography-specific factory configuration, in conjunction with Line Frequency.

**Cal. Offset** See “Checking the Fetal Recorder Offset” on page 65 and “Setting the Fetal Recorder Offset” on page 65.

## Configuring User Interface Settings

Main Setup --> User Interface

Item Name	Factory Defaults	Choice	User Defaults
QRS Volume	1	0..10	
QRS Low	0	0..10	
QRS Type <sup>1</sup>	QRS Tone	QRS Tone, QRS Tick	
Prompt Volume	8	0..10	
Tone Modulation	Yes	Yes, No	
Tone Mod. Type	Enhanced	Standard, Enhanced	
Global Speed	25mm/s	6.25, 12.5, 25, 50 mm/sec	
Touch ToneVolume	1	0..10	
Timer Volume	4	0..10	
Global SmartKeys	not applicable, these settings are stored as unique monitor settings: see "Configuring Global SmartKeys" on page 180.		
Brightness	Optimum	1..10, Optimum	
Standby Brightn.	Optimum	1..10, Optimum	
Display Units	No	Yes, No	
Alarm Limits	Yes	Yes, No	
NBP Time	Meas Time	Meas Time, Next Meas	
NBP Sys/Dia Only	Yes	Yes, No	
Wave Line Style	Thin	Thin, Medium, Thick, Extra Thick	

1.This setting can also be changed in Global settings.

## User Interface Configuration Implications

**QRS Volume** Sets the default volume of the QRS tone.

**QRS Low** Defines the minimum QRS tone volume that can be selected by the user while in Monitoring Mode.

**QRS Type** Select **QRS Tone** or **QRS Tick**. If **Tone Modulation** is set to **Yes**, the **QRS Type** automatically switches to **QRS Tone**.

**Prompt Volume** Defines the volume of the tone the monitor emits to draw the user's attention to a prompt message shown in the monitor's prompt/status line.

**Tone Modulation** if you set **Tone Modulation** to **Yes**, the pitch of the SpO<sub>2</sub> tone will change with the measured SpO<sub>2</sub> level.

**Tone Modulation Type** This setting lets you choose between **Standard** and **Enhanced**. **Standard** is the regular Nellcor behavior. **Enhanced** results in a larger (and therefore more obvious) frequency decrease for each drop in SpO<sub>2</sub> level.

**Global Speed** The **Global Speed** setting defines the speed of ECG waves on the screen.

**Touch Tone Volume** The **Touch Tone Volume** setting defines the volume of the tone you hear every time you select a field on the monitor screen. You may want to set this to 0 if you want to operate the monitor in a quiet environment.

**Timer Volume** determines the volume of the notification tone for the NST timer.

**Brightness** Defines the default brightness for monitoring.

**Standby Brightn.** Lets you choose a brightness setting for when the monitor is in Standby.

**Wave Line Style** This setting lets you configure the thickness of all waves on the screen. For better visibility over a distance you might want to use **Medium** or **Thick**. The choices are: **Thin**, **Medium**, **Thick**, **Extra Thick**.

**Alarm Limits** If **Alarm Limits** is set to **Yes**, the alarm limits are displayed next to the measurement numerics.

**NBP Time** If **NBP Time** is set to **Meas Time**, the time shown beside the NBP numeric will show the timestamp of the most recent NBP measurement. If set to **Next Meas**, and NBP mode is set to Auto, and the time until the next automatic measurement is shown.

**NBP Sys/Dia Only** If the **NBP Sys/Dia Only** setting in the User Interface menu is set to **Yes**, then only systolic and diastolic values are displayed.

## Configuring Global SmartKeys

Main Setup -> User Interface

Smartkeys			
Item Name	Factory Defaults	Complete List of SmartKeys	User Defaults
Global SmartKeys	TocoBaseline	TocoBaseline	
	FRStart/Stop	FRStart/Stop	
	PaperAdvance	PaperAdvance	
	Admit/Dischg	Admit/Dischg	
	Enter Notes	Enter Notes	
	Start/Stop (NBP)	Start/Stop (NBP)	
	Stop All (NBP)	Stop All (NBP)	
	Repeat Time (NBP)	Repeat Time (NBP)	
	Pause Alarms/Alarms Off	Pause Alarms/Alarms Off	
	Stored Data Rec	Stored Data Rec	
	Monitor Standby	Monitor Standby	
	Main Setup	Main Setup	
		Start NBP	
		Stop NBP	
		Start Recordng	
		Stop Recordng	
		Start ECG	
		Zero IUP	
		Timer	
		Set Marker	
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### Global SmartKeys Configuration Implications

**Global SmartKeys.** This lets you define the selection and sequence of the global SmartKeys.

### Changing the Selection and Sequence of Global SmartKeys

To change the selection of the Global SmartKeys:

- 1 Select **Main Setup -> User Interface -> Global SmartKeys**.
- 2 From the pop-up key line, select **Add** to open the **Choices** menu that contains all available SmartKeys.
- 3 From the **Choices** menu, select the desired SmartKey. This adds the new key to the bottom of the list of configured SmartKeys (on the left).

To delete a SmartKey from the list of configured SmartKeys,

- ◆ select it in the list, then select the pop-up key **Delete**.

To move a SmartKey to a different position,

- ◆ use the **Sort Up** and **Sort Down** pop-up keys.

# Hardware Settings

This section lists all the Hardware settings. These settings are set once per monitor. Any changes you make to the Hardware Settings configuration are automatically stored, there is no need to save them in an extra step. Hardware settings must be entered for each monitor individually, they are stored in the monitor, and they are **not cloned**.

Document the settings you configure in the empty column.

## Main Setup -> Hardware

Item Name	Factory Default	User Defaults
Calibrate Touch	n/a	
Keyboard <sup>1</sup>	US	
MIB/RS232 <sup>1</sup>	n/a	

1.Service mode only.

**Keyboard** This setting is available in **Service Mode only** and allows technical personnel to select the language of the keyboard that is connected to the P/S2 interface connector.

# Global Settings

This section lists all the Global Settings. Global Settings are set once per monitor and are independent of the User Defaults. Any changes you may configure are automatically stored, there is no need to save them.

Document the settings you configure in the empty column of the table below.

Read any information on Configuration Implications at the end of the sections before you make any configuration changes.

## Main Setup -> Global Settings

Item Name	Factory Defaults	Choice	User Defaults
Line Frequency	Geography-specific	50 Hz, 60 Hz	
QRS Type	QRS Tone	QRS Tone (most countries) QRS Tick (Japan)	
Automat. Default	Yes	Yes, No	

## Global Settings Configuration Implications

**Line Frequency** Use the **Line Frequency** setting to configure the correct line frequency for the AC Power, either 50 Hz or 60 Hz. If the Line Frequency is incorrectly set, this may affect the ECG signal quality.

**QRS Type** Select **QRS Tone** or **QRS Tick**. If **Tone Modulation** is set to **Yes**, the **QRS Type** automatically switches to **QRS Tone**.

## Automat. Default

- If **Automat. Default** is set to **Yes**, and the monitor is switched off for more than one minute, the User Defaults is reloaded in the monitor. Any unstored changes made to the active settings are lost.
- If **Automat. Default** is set to **No**, and the monitor is switched off for more than one minute, the active settings from the most recent session are retained. Automatic Default does not affect the monitor behavior when you discharge a patient. After discharge, the User Defaults is always restored.

If the monitor is switched off and then on again in less than one minute, all active settings are retained, irrespective of the **Automat. Default** setting.

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